

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:31:07 ; Search time 30 Seconds
(without alignments)
37.025 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 747574 seqs, 11073796 residues

Total number of hits satisfying chosen parameters: 135323

Minimum DB seq length: 0
Maximum DB seq length: 10

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : A_Geneseq_032802.*

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22: /SIDS1/gcgdata/geneseq/geneseq-emb1/AA2001.DAT.*
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	55	100.0	10	22 AAB46225	Human APP derived
2	52	94.5	10	22 AAB82641	All-D peptide used
3	50	90.9	9	22 AAB48493	Antifibrillogenic
4	50	90.9	10	22 AAB46224	Human APP derived
5	47	85.5	10	22 AAB46226	Human APP derived
6	46	83.6	8	18 AAW45937	Amyloid beta pepti
7	46	83.6	9	18 AAW45935	Amyloid beta pepti
8	46	83.6	10	18 AAW45934	Amyloid beta pepti
9	46	83.6	10	22 AAB46223	Human APP derived
10	42	76.4	8	17 AAW02310	Beta-amyloid modul
11	42	76.4	8	20 AAW89374	Beta-amyloid pepti

12	40	72.7	7	18	AAW45941	Amyloid beta pepti
13	40	72.7	8	18	AAW45938	Amyloid beta pepti
14	40	72.7	9	18	AAW45936	Amyloid beta pepti
15	40	72.7	10	22	AAB46222	Human APP derived
16	39	70.9	10	22	AAB46227	Human APP derived
17	38	69.1	7	17	AAW02311	Beta-amyloid prote
18	38	69.1	7	17	AAW02311	Beta-amyloid modul
19	38	69.1	7	18	AAW45940	Amyloid beta pepti
20	38	69.1	7	20	AAW89375	Beta-amyloid pepti
21	36	65.5	9	14	AAW45239	Mutant amyloid pre
22	34	61.8	6	18	AAW45946	Amyloid beta pepti
23	34	61.8	7	14	AAW45232	Beta amyloid prote
24	34	61.8	7	16	AAW87921	Test peptide used
25	34	61.8	7	16	AAW88300	Non-amnestic pepti
26	34	61.8	7	16	AAW80370	Protein polymeric
27	34	61.8	7	17	AAW02312	Beta-amyloid modul
28	34	61.8	7	17	AAW45942	Amyloid beta pepti
29	34	61.8	7	19	AAW49755	Glutamine donor pe
30	34	61.8	7	20	AAW89376	Beta-amyloid pepti
31	34	61.8	7	22	AAB67281	Residues 16-22 of
32	34	61.8	8	18	AAW45939	Amyloid beta pepti
33	34	61.8	8	18	AAW32551	Amyloidogenic sequ
34	34	61.8	8	22	AAE10663	Human amyloid prec
35	34	61.8	8	22	AAE02615	Human amyloid prec
36	34	61.8	10	21	AAW79938	Beta-amyloid recog
37	34	61.8	10	22	AAB46221	Human APP derived
38	34	61.8	10	22	AAB46228	Human APP derived
39	32	58.2	6	18	AAW45945	Amyloid beta pepti
40	31	56.4	7	14	AAW45233	Beta amyloid prote
41	31	56.4	7	22	AAB82639	All-D peptide used
42	31	56.4	7	22	AAB82640	All-D peptide used
43	31	56.4	7	22	AAB48491	Antifibrillogenic
44	31	56.4	7	22	AAB48492	Antifibrillogenic
45	30	54.5	5	18	AAW45952	Amyloid beta pepti
46	30	54.5	6	17	AAW02313	Beta-amyloid modul
47	30	54.5	6	18	AAW45947	Amyloid beta pepti
48	30	54.5	6	18	AAW45944	Amyloid beta pepti
49	30	54.5	6	20	AAW39801	Beta-amyloid prote
50	30	54.5	6	20	AAW29090	A-beta-binding pep

ALIGNMENTS

RESULT 1
AAB46225
ID AAB46225 standard; peptide; 10 AA.

XX AAB46225;

XX AC AAB46225;

XX DT 04-APR-2001 (first entry)

XX DE Human APP derived immunogenic peptide #21.

XX KW Amyloid deposit; APP; Abeta; brain; human; clearing response; neurotropic;

XX KW FC receptor mediated phagocytosis; immunogenic response; neuroprotective;

XX KW amyloid precursor protein; Alzheimer's disease.

XX OS Homo sapiens.

XX PN WO200072880-A2.

XX PD 07-DEC-2000.

XX PF 26-MAY-2000; 2000WO-US14810.

XX PR 28-MAY-1999; 99US-0322289.

XX PA (NEUR-) NEURALAB LTD.

XX PI Schenk DB, Bard F, Vasquez NJ, Yednock T;

XX WPI; 2001-032104/04.

Pub
60/00 72880
plaster 11-29-00
earlier version
by J. J. J.

XX Preventing or treating a disease associated with amyloid deposits,
PT especially Alzheimer's disease, comprises administering amyloid
PT specific antibody -
XX
PS Disclosure; Figure 19; 143pp; English.
XX
CC This invention describes a novel method of preventing or treating a
CC disease associated with amyloid deposits of amyloid precursor protein
CC (APP) Abeta fragments in the brain of a patient, which comprises
CC administering to the patient: (a) an antibody that binds to Abeta, the
CC antibody binds to an amyloid deposit and induces a clearing response (Fc
CC receptor mediated phagocytosis) against it (b) a polypeptide containing
CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
CC that induces an immunogenic response against residues 1-3 to 7-11 of
CC Abeta. The products of the invention have neurotropic and neuroprotective
CC activity. The method is also useful for monitoring a course of treatment
CC being administered to a patient e.g. active and passive immunization. The
CC methods are useful for prophylactic and therapeutic treatment of
CC Alzheimer's disease.
XX
SQ Sequence 10 AA;
Query Match 100.0%; Score 55; DB 22; Length 10;
Best Local Similarity 100.0%; Pred. No. 0.00014; Mismatches 0; Indels 0; Gaps 0;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
DB 1 HHQKLVFFAE 10
|||||||
RESULT 2
ID AAB82641 standard; Peptide; 10 AA.
XX
AC AAB82641;
XX
DT 02-OCT-2001 (first entry)
XX
DE All-D peptide used in Alzheimer's disease vaccine.
XX
KW Alzheimer's disease; amyloidosis; amyloid-related disease;
KW vaccine; therapy; antigen.
XX
OS Synthetic.
XX
FH Key Location/Qualifiers
FT Misc-difference 1..10 /note= "all D-form residues"
XX
PN WO200139796-A2.
XX
PD 07-JUN-2001.
XX
PF 29-NOV-2000; 2000WO-CA01413.
XX
PR 29-NOV-1999; 99US-0168594.
XX
PR 28-NOV-2000; 2000US-0724842.
XX
PA (NEUR-) NEUROCHEM INC.
XX
PI Chalfour R, Hebert L, Kong X, Gervais F;
XX
DR WPI; 2001-441458/47.
XX
PT Preventing/treating amyloid-related disease, especially Alzheimer's
PT disease, comprises administering antigenic all-D peptide, e.g. as
PT vaccine, which elicits production of antibodies to prevent
PT fibrillogenesis and associated cellular toxicity -
XX
PS Disclosure; Page 11; 31pp; English.
XX

CC The present sequence is that of an all-D peptide suitable for
CC use in preparing vaccines for preventing or treating Alzheimer's
CC disease and other amyloid related disorders in humans. It is based
CC on a portion of amyloid-beta peptide (see AAB82622), and may be
CC modified by removing or inserting 1 or more amino acid residues, or
CC by substituting 1 or more amino acid residues with other amino acid
CC residues or non-amino acid fragments. Vaccines of the invention
CC are produced using 'non-self' peptides synthesised from the
CC unnatural D-configuration amino acids to avoid the drawbacks of
CC 'self' proteins. The all-D peptides need not be aggregated to be
CC operative or immunogenic. They preferably interact with at
CC least 1 region of an amyloid protein, e.g. the beta-sheet region
CC or GAG-binding site region, the amyloid-beta peptide, or their
CC immunogenic fragments, protein conjugates, immunogenic derivative
CC peptides and immunogenic peptidomimetics. Examples include all-D
CC peptides corresponding to residues 1-42, 1-40, 1-35, 1-28, 1-7,
CC 10-16, 16-21 and 36-42 of the amyloid-beta peptide and the all-D
CC derivative peptides given in AAB82623-64. The vaccine elicits a
CC preferential TH-2 or TH-1 response, preventing fibrillogenesis and
CC associated cellular toxicity. The amyloid related diseases may be
CC localised amyloidosis, e.g. diabetes type II, neurodegenerative
CC diseases, e.g. bovine spongiform encephalitis, Creutzfeldt-Jakob
CC disease, scrapie, cerebral amyloid angiopathy, and prion protein
CC related disorders, or systemic amyloidosis associated with chronic
CC infection (e.g. tuberculosis) or chronic inflammation (e.g.
CC rheumatoid arthritis), familial Mediterranean fever (FMF) and
CC systemic amyloidosis found in long-term haemodialysis patients.
XX
SQ Sequence 10 AA;
Query Match 94.5%; Score 52; DB 22; Length 10;
Best Local Similarity 90.0%; Pred. No. 0.00056; Mismatches 1; Indels 0; Gaps 0;
Matches 9; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
DB 1 HHQKLVFFAQ 10
|||||||
RESULT 3
ID AAB48493 standard; Peptide; 9 AA.
XX
AC AAB48493;
XX
DT 02-MAR-2001 (first entry)
XX
DE Antifibrillogenic peptide #20.
XX
KW Nootropic; neuroprotective; antifibrillogenic; amyloidosis inhibition;
KW cytoprotection; amyloid deposit degradation; amyloidosis disorder;
KW Alzheimer's disease.
XX
OS Homo sapiens.
XX
FH Key Location/Qualifiers
FT Modified-site 9 /note= "C-terminal amide"
XX
PN WO200068263-A2.
XX
PD 16-NOV-2000.
XX
PF 04-MAY-2000; 2000WO-CA00515.
XX
PR 05-MAY-1999; 99US-0132592.
XX
PA (NEUR-) NEUROCHEM INC.
XX
PI Chalfour R, Gervais F, Gupta A;
XX
DR WPI; 2001-031852/04.
XX

PT Antifibrillogenic agent useful for inhibiting amyloidosis and/or for
 PT cytoprotection for treating amyloidosis disorders, comprises a peptide,
 PT its isomer or peptidomimetic -
 XX
 PS Claim 7; Page 25; 46pp; English.
 XX
 CC Peptides AAB48474-B48496 are antifibrillogenic agents that can be used
 CC for inhibiting amyloidosis and/or for cytoprotection. The peptides of
 CC AAB48474-B48496 cause the breakdown of amyloid deposits and are
 CC therefore useful for treating amyloidosis disorders such as Alzheimer's
 CC disease. Peptides AAB48474-B48496 were identified from the
 CC glycosaminoglycan binding region and the prot-prot interaction region of
 CC the human amyloid protein.
 XX

SQ Sequence 9 AA;
 Query Match 90.9%; Score 50; DB 22; Length 9;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFFA 9
 Db 1 HHQKLVFFFA 9

RESULT 4

AAB46224
 ID AAB46224 standard; peptide; 10 AA.

XX AAB46224;

XX 04-APR-2001 (first entry)

DE Human APP derived immunogenic peptide #20.

XX Amyloid deposit; APP; Abeta; brain; human; clearing response; nootropic;
 KW Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
 KW amyloid precursor protein; Alzheimer's disease.

XX Homo sapiens.

XX WO200072880-A2.

XX 07-DEC-2000.

XX 26-MAY-2000; 2000WO-US14810.

XX 28-MAY-1999; 99US-0322289.

XX (NEUR-) NEURALAB LTD.

XX Schenk DB, Bard F, Vasquez NJ, Yednock T;

XX WPI; 2001-032104/04.

XX Preventing or treating a disease associated with amyloid deposits,
 PT especially Alzheimer's disease, comprises administering amyloid
 PT specific antibody -

XX Disclosure; Figure 19; 143pp; English.
 XX

CC This invention describes a novel method of preventing or treating a
 CC disease associated with amyloid deposits of amyloid precursor protein
 CC (APP) Abeta fragments in the brain of a patient, which comprises
 CC administering to the patient: (a) an antibody that binds to Abeta, the
 CC antibody binds to an amyloid deposit and induces a clearing response (Fc
 CC receptor mediated phagocytosis) against it (b) a polypeptide containing
 CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
 CC that induces an immunogenic response against residues 1-3 to 7-11 of
 CC Abeta. The products of the invention have nootropic and neuroprotective
 CC activity. The method is also useful for monitoring a course of treatment
 CC being administered to a patient e.g. active and passive immunization. The
 CC methods are useful for prophylactic and therapeutic treatment of

CC Alzheimer's disease.

XX Sequence 10 AA;

Query Match 90.9%; Score 50; DB 22; Length 10;
 Best Local Similarity 100.0%; Pred. No. 0.0014;
 Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFFA 9

Db 2 HHQKLVFFFA 10

RESULT 5

AAB46226
 ID AAB46226 standard; peptide; 10 AA.

XX AAB46226;

XX 04-APR-2001 (first entry)

XX Human APP derived immunogenic peptide #22.

XX Amyloid deposit; APP; Abeta; brain; human; clearing response; nootropic;
 KW Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
 KW amyloid precursor protein; Alzheimer's disease.

XX Homo sapiens.

XX WO200072880-A2.

XX 07-DEC-2000.

XX 26-MAY-2000; 2000WO-US14810.

XX 28-MAY-1999; 99US-0322289.

XX (NEUR-) NEURALAB LTD.

XX Schenk DB, Bard F, Vasquez NJ, Yednock T;

XX WPI; 2001-032104/04.

XX Preventing or treating a disease associated with amyloid deposits,
 PT especially Alzheimer's disease, comprises administering amyloid
 PT specific antibody -

XX Disclosure; Figure 19; 143pp; English.
 XX

CC This invention describes a novel method of preventing or treating a
 CC disease associated with amyloid deposits of amyloid precursor protein
 CC (APP) Abeta fragments in the brain of a patient, which comprises
 CC administering to the patient: (a) an antibody that binds to Abeta, the
 CC antibody binds to an amyloid deposit and induces a clearing response (Fc
 CC receptor mediated phagocytosis) against it (b) a polypeptide containing
 CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
 CC that induces an immunogenic response against residues 1-3 to 7-11 of
 CC Abeta. The products of the invention have nootropic and neuroprotective
 CC activity. The method is also useful for monitoring a course of treatment
 CC being administered to a patient e.g. active and passive immunization. The
 CC methods are useful for prophylactic and therapeutic treatment of

XX Sequence 10 AA;

Query Match 85.5%; Score 47; DB 22; Length 10;
 Best Local Similarity 100.0%; Pred. No. 0.0054;
 Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFFAE 10

Db 1 HOKLVFFFAE 9

RESULT 6
AAW45937
ID AAW45937 standard; peptide; 8 AA.
XX
AC AAW45937;
XX
DT 30-JUN-1998 (first entry)
XX
DE Amyloid beta peptide fragment.
XX
DE Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.
XX
OS Homo sapiens.
XX
PN WO9721728-A1.
XX
PD 19-JUN-1997.
XX
PF 09-DEC-1996; 96WO-SE01621.
XX
PR 29-DEC-1995; 95US-0009386.
XX
PR 12-DEC-1995; 95SE-0004467.
XX
PA (KARO-) KAROLINSKA INNOVATIONS AB.
XX
PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
XX
PD WPI; 1997-332723/30.
XX
PF 09-DEC-1996; 96WO-SE01621.
XX
PR 29-DEC-1995; 95US-0009386.
XX
PR 12-DEC-1995; 95SE-0004467.
XX
PA (KARO-) KAROLINSKA INNOVATIONS AB.
XX
PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
XX
PD WPI; 1997-332723/30.
XX
PF Use of new and known peptide(s) for inhibition of polymerisation of
PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
PT Down's syndrome associated with amyloidosis.
XX
PS Example 1; Figure 2B; 31pp; English.
XX
CC This sequence represents a fragment of the amyloid beta peptide. The
CC invention relates to the use of peptide compounds for inhibition of
CC polymerisation of amyloid beta peptide (ABP), as model substances for
CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
CC tool for the identification of other organic compounds with similar
CC functional properties, or as ligands in positron emission tomography.
CC The peptides may be used in treatment of amyloidosis, especially in
CC treatment of Alzheimer's disease associated with amyloidosis, for
CC treatment or prevention of demens in patients with Down's syndrome, for
CC treatment or prevention of hereditary cerebral haemorrhage with
CC amyloidosis (Dutch type) or for the prevention of fibril formation of
CC human amyloid protein. They can also be used for identifying other
CC molecules with similar properties and/or as ligands for detection of
CC amyloid deposits using e.g. positron emission tomography.
XX
SQ Sequence 8 AA;
Query Match 83.6%; Score 46; DB 18; Length 8;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFF 8
DB 1 HHQKLVFF 8
RESULT 7
AAW45935
ID AAW45935 standard; peptide; 9 AA.
XX
AC AAW45935;
XX
DT 08-JUL-1998 (first entry)
XX
DE Amyloid beta peptide fragment.

KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.
XX
OS Homo sapiens.
XX
PN WO9721728-A1.
XX
PD 19-JUN-1997.
XX
PF 09-DEC-1996; 96WO-SE01621.
XX
PR 29-DEC-1995; 95US-0009386.
XX
PR 12-DEC-1995; 95SE-0004467.
XX
PA (KARO-) KAROLINSKA INNOVATIONS AB.
XX
PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
XX
PD WPI; 1997-332723/30.
XX
PF Use of new and known peptide(s) for inhibition of polymerisation of
PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
PT Down's syndrome associated with amyloidosis.
XX
PS Example 1; Figure 2B; 31pp; English.
XX
CC This sequence represents a fragment of the amyloid beta peptide. The
CC invention relates to the use of peptide compounds for inhibition of
CC polymerisation of amyloid beta peptide (ABP), as model substances for
CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
CC tool for the identification of other organic compounds with similar
CC functional properties, or as ligands in positron emission tomography.
CC The peptides may be used in treatment of amyloidosis, especially in
CC treatment of Alzheimer's disease associated with amyloidosis, for
CC treatment or prevention of demens in patients with Down's syndrome, for
CC treatment or prevention of hereditary cerebral haemorrhage with
CC amyloidosis (Dutch type) or for the prevention of fibril formation of
CC human amyloid protein. They can also be used for identifying other
CC molecules with similar properties and/or as ligands for detection of
CC amyloid deposits using e.g. positron emission tomography.
XX
SQ Sequence 9 AA;
Query Match 83.6%; Score 46; DB 18; Length 9;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFF 8
DB 2 HHQKLVFF 9
RESULT 8
AAW45934
ID AAW45934 standard; peptide; 10 AA.
XX
AC AAW45934;
XX
DT 08-JUL-1998 (first entry)
XX
DE Amyloid beta peptide fragment (residues 11-20).
XX
KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.
XX
OS Homo sapiens.
XX
PN WO9721728-A1.
XX
PD 19-JUN-1997.
XX
PF 09-DEC-1996; 96WO-SE01621.
XX

PR 29-DEC-1995; 95US-0009386.
 PR 12-DEC-1995; 95SE-0004467.
 XX
 PA (KARO-) KAROLINSKA INNOVATIONS AB.
 XX
 PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 XX
 DR WPI; 1997-332723/30.
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 XX
 PS Example 1: Page 10; 31pp; English.
 XX
 CC This sequence represents a fragment of the amyloid beta peptide
 CC (residues 11-20). The invention relates to the use of peptide
 CC compounds for inhibition of polymerisation of amyloid beta peptide
 CC (ABP), as model substances for synthesis of ABP-ligands for inhibition
 CC of polymerisation of ABP, as a tool for the identification of other
 CC organic compounds with similar functional properties, or as ligands in
 CC positron emission tomography. The peptides may be used in treatment of
 CC amyloidosis, especially in treatment of Alzheimer's disease associated
 CC with amyloidosis, for treatment or prevention of demens in patients with
 CC Down's syndrome, for treatment or prevention of hereditary cerebral
 CC haemorrhage with amyloidosis (Dutch type) or for the prevention of
 CC fibril formation of human amyloid protein. They can also be used for
 CC identifying other molecules with similar properties and/or as ligands
 CC for detection of amyloid deposits using e.g. positron emission
 CC tomography.
 XX
 SQ Sequence 10 AA;
 Query Match 83.6%; Score 46; DB 18; Length 10;
 Best Local Similarity 100.0%; Pred. No. 0.0086;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Qy 1 HHQKLVFF 8
 Db 3 HHQKLVFF 10
 |||||
 RESULT 9
 AAB46223
 ID AAB46223 standard; peptide; 10 AA.
 XX
 AC AAB46223;
 XX
 DT 04-APR-2001 (first entry)
 XX
 DE Human APP derived immunogenic peptide #19.
 XX
 KW Amyloid deposit; APP; Abeta; brain; human; clearing response; neurotropic;
 KW Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
 KW amyloid precursor protein; Alzheimer's disease.
 XX
 OS Homo sapiens.
 XX
 FN WO200072880-A2.
 XX
 XX 07-DEC-2000.
 PD
 XX
 PF 26-MAY-2000; 2000WO-US14810.
 XX
 XX 28-MAY-1999; 99US-0322289.
 PR
 XX (NEUR-) NEURALAB LTD.
 PA
 XX
 XX Schenk DB, Bard F, Vasquez NJ, Yednock T;
 PI
 XX WPI; 2001-032104/04.
 DR
 XX Preventing or treating a disease associated with amyloid deposits,
 PT

PT especially Alzheimer's disease, comprises administering amyloid
 PT specific antibody -
 XX
 PS Disclosure; Figure 19; 143pp; English.
 XX
 CC This invention describes a novel method of preventing or treating a
 CC disease associated with amyloid deposits of amyloid precursor protein
 CC (APP) Abeta fragments in the brain of a patient, which comprises
 CC administering to the patient: (a) an antibody that binds to Abeta, the
 CC antibody binds to an amyloid deposit and induces a clearing response (Fc
 CC receptor mediated phagocytosis) against it (b) a polypeptide containing
 CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
 CC that induces an immunogenic response against residues 1-3 to 7-11 of
 CC Abeta. The products of the invention have neurotropic and neuroprotective
 CC activity. The method is also useful for monitoring a course of treatment
 CC being administered to a patient e.g. active and passive immunization. The
 CC methods are useful for prophylactic and therapeutic treatment of
 CC Alzheimer's disease.
 XX
 SQ Sequence 10 AA;
 Query Match 83.6%; Score 46; DB 22; Length 10;
 Best Local Similarity 100.0%; Pred. No. 0.0086;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Qy 1 HHQKLVFF 8
 Db 3 HHQKLVFF 10
 |||||
 RESULT 10
 AAW02310
 ID AAW02310 standard; peptide; 8 AA.
 XX
 AC AAW02310;
 XX
 DT 02-MAY-1997 (first entry)
 XX
 DE Beta-amyloid modulator peptide #1.
 XX
 KW Beta-amyloid; modulator; amyloid plaque; brain lesion; amyloidosis;
 KW cerebral blood vessel; Alzheimer's disease; amyloidogenic protein;
 KW familial amyloid polyneuropathy; familial amyloid cardiomyopathy;
 KW isolated cardiac amyloidosis; systemic senile amyloidosis; insulinoma;
 KW bovine spongiform encephalopathy; Creutzfeldt-Jakob disease; urticaria;
 KW adult-onset diabetes; familial Mediterranean fever; therapy; deafness;
 KW scrapie; familial amyloid nephropathy; hereditary cerebral haemorrhage.
 XX
 OS Synthetic.
 XX
 XX WO9628471-A1.
 PN
 XX 19-SEP-1996.
 PD
 XX 14-MAR-1996; 96WO-US03492.
 PF
 XX 27-OCT-1995; 95US-0548998.
 PR
 XX 14-MAR-1995; 95US-0404831.
 PR
 XX 07-JUN-1995; 95US-0475579.
 PR
 XX (PHAR-) PHARM PEPTIDES INC.
 PA
 XX Benjamin H, Chin J, Findeis MA, Garnick MB, Geffer ML;
 PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
 PI Molineaux S, Musso G, Reed MJ, Signer ER, Wakefield J;
 XX
 XX WPI; 1996-433762/43.
 DR
 XX Modulators of amyloid aggregation - comprising, e.g. amyloidogenic
 PT protein coupled (indirectly) to at least 1 modifying gp., useful in
 PT treatment of Alzheimer's disease
 XX
 PS Claim 16; Page 90; 106pp; English.

XX AAW02310-W02332 represent the peptide portions of the beta-amyloid
 CC modulator compounds of the invention. Beta-amyloid peptide is a 4
 CC kilodalton peptide that is the major protein component of amyloid
 CC plaques. Amyloid plaques are present both in the brain lesions, and in
 CC the walls of cerebral blood vessels in Alzheimer's disease patients.
 CC The amyloid modulators of the invention comprise an amyloidogenic protein
 CC or peptide (such as this sequence) coupled directly or indirectly to at
 CC least one modifying group. The modifying group is preferably a cyclic,
 CC heterocyclic, or polycyclic group, such as decalin, a cholanyl group, a
 CC biotin containing group, or a fluorescein containing group. These
 CC compounds then modulate the aggregation of these sequences to natural
 CC amyloid proteins or peptides when contacted with the natural
 CC amyloidogenic proteins or peptides. The modulator compounds can be used
 CC in the treatment of disorders associated with amyloidosis, such as
 CC familial amyloid polynuropathy, familial amyloid cardiomyopathy,
 CC isolated cardiac amyloidosis, systemic senile amyloidosis, scrapie,
 CC bovine spongiform encephalopathy, Creutzfeldt-Jakob disease, adult-onset
 CC diabetes, insulinoma, familial Mediterranean fever, familial amyloid
 CC nephropathy with urticaria and deafness, hereditary cerebral haemorrhage
 CC and other types of amyloidosis. The modulators are also useful for the
 CC treatment of disorders associated with beta-amyloidosis, especially
 CC Alzheimer's disease.
 XX
 SQ Sequence 8 AA;

Query Match 76.4%; Score 42; DB 17; Length 8;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFA 9
 |||||
 Db 1 HOKLVFFA 8

RESULT 11
 AAW89374
 ID AAW89374 standard; peptide; 8 AA.

AC AAW89374;

DT 02-MAR-1999 (first entry)

DE Beta-amyloid peptide derivative A-beta-14-21.

XX Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
 KW aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
 KW familial amyloid polynuropathy; bovine spongiform encephalopathy;
 KW Creutzfeldt-Jakob disease; BAP.

XX Homo sapiens.
 OS Synthetic.

XX US5854204-A.

XX 29-DEC-1998.

XX 14-MAR-1996; 96US-0612785.

XX 14-MAR-1996; 96US-0612785.

PR 14-MAR-1995; 95US-0404831.

PR 07-JUN-1995; 95US-0475579.

PR 27-OCT-1995; 95US-0548998.

XX (PRAE-) PRAECIS PHARM INC.

XX Benjamin H, Chin J, Findeis WA, Garnick WB, Geffer ML;

PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;

PI Mollineaux S, Musso G, Reed M, Signer ER, Wakefield J;

XX WPI; 1999-094964/08.

PT New peptide(s) derived from beta-amyloid peptide that inhibit

PT amyloid aggregation - and neurotoxicity, specifically for treatment
 PT and prevention of Alzheimer's disease
 XX
 XX Example 12; Column 64; 52pp; English.

CC The present invention describes beta-amyloid peptide (BAP) derivatives.
 CC The BAP derivatives inhibit aggregation of amyloidogenic proteins and
 CC peptides, specifically BAP, and their neurotoxicity, so are useful for
 CC treating and preventing any disease involving amyloidosis, specifically
 CC Alzheimer's disease but also Down's syndrome, familial amyloid
 CC polynuropathy or cardiomyopathy, bovine spongiform encephalopathy and
 CC Creutzfeldt-Jakob disease. The BAP derivatives are also used to diagnose
 CC these diseases, in vitro or in vivo, by detecting binding of BAP to
 CC labelled BAP derivatives. Some BAP derivatives inhibit BAP aggregation
 CC even when BAP is present in molar excess. The present sequence
 CC represents a BAP derivative.

SQ Sequence 8 AA;

Query Match 76.4%; Score 42; DB 20; Length 8;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFA 9
 |||||
 Db 1 HOKLVFFA 8

RESULT 12
 AAW45941
 ID AAW45941 standard; peptide; 7 AA.

XX AAW45941;

XX 30-JUN-1998 (first entry)

XX Amyloid beta peptide fragment.

XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

XX WO9721728-A1.

XX 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

PR 29-DEC-1995; 95US-0009386.

PR 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

XX WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.

XX Example 1; Figure 2B; 31pp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of demens in patients with Down's syndrome, for

CC treatment or prevention of hereditary cerebral haemorrhage with
CC amyloidosis (Dutch type) or for the prevention of fibril formation of
CC human amyloid protein. They can also be used for identifying other
CC molecules with similar properties and/or as ligands for detection of
CC amyloid deposits using e.g. positron emission tomography.
XX
SQ Sequence 7 AA;

Query Match 72.7%; Score 40; DB 18; Length 7;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVF 7
| | | | |
Db 1 HHQKLVF 7

RESULT 13
AAW45938
ID AAW45938 standard; peptide; 8 AA.

XX AAW45938;

AC AAW45938;

DT 30-JUN-1998 (first entry)

DE Amyloid beta peptide fragment.

KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

PN WO9721728-A1.

PD 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

XX 29-DEC-1995; 95US-0009386.

PR 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

XX WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of

PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or

PT Down's syndrome associated with amyloidosis.

XX Example 1; Figure 2B; 3lpp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The

CC invention relates to the use of peptide compounds for inhibition of

CC polymerisation of amyloid beta peptide (ABP), as model substances for

CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a

CC tool for the identification of other organic compounds with similar

CC functional properties, or as ligands in positron emission tomography.

CC The peptides may be used in treatment of amyloidosis, especially in

CC treatment of Alzheimer's disease associated with amyloidosis, for

CC treatment or prevention of demens in patients with Down's syndrome, for

CC treatment or prevention of hereditary cerebral haemorrhage with

CC amyloidosis (Dutch type) or for the prevention of fibril formation of

CC human amyloid protein. They can also be used for identifying other

QY 1 HHQKLVF 7
| | | | |
Db 2 HHQKLVF 8

RESULT 14
AAW45936

ID AAW45936 standard; peptide; 9 AA.

XX AAW45936;

XX 30-JUN-1998 (first entry)

XX Amyloid beta peptide fragment.

XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

PN WO9721728-A1.

XX 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

XX 29-DEC-1995; 95US-0009386.

PR 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

XX WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of

PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or

PT Down's syndrome associated with amyloidosis.

XX Example 1; Figure 2B; 3lpp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The

CC invention relates to the use of peptide compounds for inhibition of

CC polymerisation of amyloid beta peptide (ABP), as model substances for

CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a

CC tool for the identification of other organic compounds with similar

CC functional properties, or as ligands in positron emission tomography.

CC The peptides may be used in treatment of amyloidosis, especially in

CC treatment of Alzheimer's disease associated with amyloidosis, for

CC treatment or prevention of demens in patients with Down's syndrome, for

CC treatment or prevention of hereditary cerebral haemorrhage with

CC amyloidosis (Dutch type) or for the prevention of fibril formation of

CC human amyloid protein. They can also be used for identifying other

CC molecules with similar properties and/or as ligands for detection of

CC amyloid deposits using e.g. positron emission tomography.

XX Sequence 9 AA;

Query Match 72.7%; Score 40; DB 18; Length 9;

Best Local Similarity 100.0%; Pred. No. 6.4e+05;

Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVF 7
| | | | |
Db 3 HHQKLVF 9

RESULT 15
AAB46222

ID AAB46222 standard; peptide; 10 AA.

XX AAB46222;

AC AAB46222;

XX New mutant forms of amyloid precursor protein - for detecting
PT cpds. that modify activity of enzymes involved in precursor
PT cleavage, also new nucleic acid encoding them
XX
PS Disclosure; Page 34; 66pp; English.
XX
CC Recombinant polypeptides produced using the coding sequences of
CC mutant forms of amyloid precursor proteins comprising from the 5' to
CC the 3' end a sequence encoding a marker and either (1) a sequence
CC encoding the N-terminus of an amyloid precursor protein (APP) up to,
CC but not including, the nucleotides encoding the beta amyloid protein
CC (BAP) domain or (2) the BAP domain; or the two ligated together, can
CC be used to detect drugs or compounds that inhibit/augment the
CC activity of proteolytic enzymes which cleave APP to generate BAP
CC fragments (deposition of which occurs in patients with Alzheimers
CC disease and Down's syndrome). This fragment corresponding to amino
CC acid residues 14-20 of BAP can be altered and affect the level of
CC secretion of APP's containing the BAP sequence.
XX
SQ Sequence 7 AA;
Query Match 69.1%; Score 38; DB 14; Length 7;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Oy 2 HQKLVFF 8
Db 1 HQKLVFF 7
RESULT 18
AAW02311
ID AAW02311 standard; peptide; 7 AA.
XX
AC AAW02311;
XX
DT 02-MAY-1997 (first entry)
XX
DE Beta-amyloid modulator peptide #2.
XX
KW Beta-amyloid; modulator; amyloid plaque; brain lesion; amyloidosis;
KW cerebral blood vessel; Alzheimer's disease; amyloidogenic protein;
KW familial amyloid polyneuropathy; familial amyloid cardiomyopathy;
KW isolated cardiac amyloidosis; systemic senile amyloidosis; insulinoma;
KW bovine spongiform encephalopathy; Creutzfeldt-Jakob disease; urticaria;
KW adult-onset diabetes; familial Mediterranean fever; therapy; deafness;
KW scrapie; familial amyloid nephropathy; hereditary cerebral haemorrhage.
XX
OS Synthetic.
XX
PN W09628471-A1.
XX
PD 19-SEP-1996.
XX
PF 14-MAR-1996; 96WO-US03492.
XX
PR 27-OCT-1995; 95US-0548998.
XX
PR 14-MAR-1995; 95US-0404831.
XX
PR 07-JUN-1995; 95US-0475579.
XX
PA (PHAR-) PHARM PEPTIDES INC.
XX
PI Benjamin H, Chin J, Findeis MA, Garnick MB, Geftel ML;
PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
PI Molineaux S, Musso G, Reed MJ, Signer ER, Wakefield J;
XX
DR WPI; 1996-433762/43.
XX
XX Modulators of amyloid aggregation - comprising, e.g. amyloidogenic
PT protein coupled (indirectly to at least 1 modifying gp., useful in
PT treatment of Alzheimer's disease
XX

PS Claim 16; Page 90; 106pp; English.
XX
CC AAW02310-W02332 represent the peptide portions of the beta-amyloid
CC modulator compounds of the invention. Beta-amyloid peptide is a 4
CC kilodalton peptide that is the major protein component of amyloid
CC plaques. Amyloid plaques are present both in the brain lesions, and in
CC the walls of cerebral blood vessels in Alzheimer's disease patients.
CC The amyloid modulators of the invention comprise an amyloidogenic protein
CC or peptide (such as this sequence) coupled directly or indirectly to at
CC least one modifying group. The modifying group is preferably a cyclic,
CC heterocyclic, or polycyclic group, such as decalin, a cholanyl group, a
CC biotin containing group, or a fluorescein containing group. These
CC compounds then modulate the aggregation of these sequences to natural
CC amyloid proteins or peptides when contacted with the natural
CC amyloidogenic proteins or peptides. The modulator compounds can be used
CC in the treatment of disorders associated with amyloidosis, such as
CC familial amyloid polyneuropathy, familial amyloid cardiomyopathy,
CC isolated cardiac amyloidosis, systemic senile amyloidosis, scrapie,
CC bovine spongiform encephalopathy, Creutzfeldt-Jakob disease, adult-onset
CC diabetes, insulinoma, familial Mediterranean fever, familial amyloid
CC nephropathy with urticaria and deafness, hereditary cerebral haemorrhage
CC and other types of amyloidosis. The modulators are also useful for the
CC treatment of disorders associated with beta-amyloidosis, especially
CC Alzheimer's disease.
XX
SQ Sequence 7 AA;
Query Match 69.1%; Score 38; DB 17; Length 7;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Oy 2 HQKLVFF 8
Db 1 HQKLVFF 7
RESULT 19
AAW45940
ID AAW45940 standard; peptide; 7 AA.
XX
AC AAW45940;
XX
DT 30-JUN-1998 (first entry)
XX
DE Amyloid beta peptide fragment.
XX
KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.
XX
OS Homo sapiens.
XX
PN W09721728-A1.
XX
PD 19-JUN-1997.
XX
PF 09-DEC-1996; 96WO-SE01621.
XX
PR 29-DEC-1995; 95US-0009386.
XX
PR 12-DEC-1995; 95SE-0004467.
XX
PA (KARO-) KAROLINSKA INNOVATIONS AB.
XX
PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
XX WPI; 1997-332723/30.
XX
XX Use of new and known peptide(s) for inhibition of polymerisation of
PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
PT Down's syndrome associated with amyloidosis.
XX
XX Example 1; Figure 2B; 31pp; English.
XX
XX This sequence represents a fragment of the amyloid beta peptide. The

CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 7 AA;
 Query Match 59.1%; Score 38; DB 18; Length 7;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2 HQKLVEFF 8
 Db 1 HQKLVEFF 7
 RESULT 20
 AAW89375
 ID AAW89375 standard; peptide; 7 AA.
 AC AAW89375;
 XX
 DT 02-MAR-1999 (first entry)
 DE Beta-amyloid peptide derivative A-beta-14-20.
 XX
 KW Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
 KW aggregation, neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
 KW familial amyloid polynuropathy; bovine spongiform encephalopathy;
 KW Creutzfeldt-Jakob disease; BAP.
 XX
 OS Homo sapiens.
 OS Synthetic.
 XX
 PN US5854204-A.
 XX
 PD 29-DEC-1998.
 XX
 PF 14-MAR-1996; 96US-0612785.
 XX
 PR 14-MAR-1996; 96US-0612785.
 PR 14-MAR-1995; 95US-0404831.
 PR 07-JUN-1995; 95US-0475579.
 PR 27-OCT-1995; 95US-0548998.
 XX
 PA (PRAE-) PRAECIS PHARM INC.
 XX
 PI Benjamin H, Chin J, Findels MA, Garnick MB, Gefter ML;
 PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
 PI Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;
 XX
 DR WPI; 1999-094964/08.
 XX
 PT New peptide(s) derived from beta-amyloid peptide that inhibit
 PT amyloid aggregation - and neurotoxicity, specifically for treatment
 PT and prevention of Alzheimer's disease
 XX
 PS Example 12; Column 64; 52pp; English.
 XX
 CC The present invention describes beta-amyloid peptide (BAP) derivatives.
 CC The BAP derivatives inhibit aggregation of amyloidogenic proteins and
 CC peptides, specifically BAP, and their neurotoxicity, so are useful for
 CC treating and preventing any disease involving amyloidosis, specifically
 CC Alzheimer's disease but also Down's syndrome, familial amyloid

CC polynuropathy or cardiomyopathy, bovine spongiform encephalopathy and
 CC Creutzfeldt-Jakob disease. The BAP derivatives are also used to diagnose
 CC these diseases, in vitro or in vivo, by detecting binding of BAP to
 CC labelled BAP derivatives. Some BAP derivatives inhibit BAP aggregation
 CC even when BAP is present in molar excess. The present sequence
 CC represents a BAP derivative.
 XX
 SQ Sequence 7 AA;
 Query Match 59.1%; Score 38; DB 20; Length 7;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2 HQKLVEFF 8
 Db 1 HQKLVEFF 7
 RESULT 21
 AAR45239
 ID AAR45239 standard; Peptide; 9 AA.
 XX
 AC AAR45239;
 XX
 DT 20-JUN-1994 (first entry)
 DE Mutant amyloid precursor protein fragment.
 XX
 KW Amyloid precursor protein; APP; beta amyloid protein; BAP;
 KW detection; Alzheimer's disease; Down's syndrome.
 XX
 OS Homo sapiens.
 XX
 PN AU9338358-A.
 XX
 PD 04-NOV-1993.
 XX
 PF 03-MAY-1993; 93AU-0038358.
 XX
 PR 01-MAY-1992; 92US-0877675.
 XX
 PA (AMCY) AMERICAN CYANAMID CO.
 XX
 PI Jacobsen JS, Vitek MP;
 XX
 DR WPI; 1993-406194/51.
 DR N-PSDB; AAQ54267.
 XX
 PT New mutant forms of amyloid precursor protein - for detecting
 PT cpds. that modify activity of enzymes involved in precursor
 PT cleavage, also new nucleic acid encoding them
 XX
 PS Disclosure; Page 35; 66pp; English.
 XX
 CC Recombinant polypeptides produced using the coding sequences of
 CC mutant forms of amyloid precursor proteins comprising from the 5' to
 CC the 3' end a sequence encoding a marker and either (1) a sequence
 CC encoding the N-terminus of an amyloid precursor protein (APP) up to,
 CC but not including, the nucleotides encoding the beta amyloid protein
 CC (BAP) domain or (2) the BAP domain; or the two ligated together, can
 CC be used to detect drugs or compounds that inhibit/augment the
 CC activity of proteolytic enzymes which cleave APP to generate BAP
 CC fragments (deposition of which occurs in patients with Alzhemiers
 CC disease and Down's syndrome). This is a fragment of amyloid
 CC precursor protein containing a mutation which is associated with
 CC diseases involving BAP deposition.
 XX
 SQ Sequence 9 AA;
 Query Match 65.5%; Score 36; DB 14; Length 9;
 Best Local Similarity 87.5%; Pred. No. 6.4e+05;
 Matches 7; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 3 OKLVEFAE 10
 |||||
 Db 1 OKLVEFAQ 8

RESULT 22

AAW45946
 ID AAW45946 standard; peptide; 6 AA.

AC AAW45946;

XX 30-JUN-1998 (first entry)

XX Amyloid beta peptide fragment.

XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

PN WO9721728-Al.

PD 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

XX 29-DEC-1995; 95US-0009386.

PR 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

DR WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.

PS Example 1; Figure 2B; 31pp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.

XX Sequence 6 AA;

Query Match 61.8%; Score 34; DB 18; Length 6;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6

Db 1 HHQKLV 6

RESULT 23

AAW45232

ID AAR45232 standard; Peptide; 7 AA.

XX AAR45232;

AC AAR45232;

XX

DT 20-JUN-1994 (first entry)

XX Beta amyloid protein fragment.

XX Amyloid precursor protein; APP; beta amyloid protein; BAP;
 KW detection; Alzheimer's disease; Down's syndrome.

XX Homo sapiens.

PN AU9338358-A.

XX 04-NOV-1993.

XX 03-MAY-1993; 93AU-0038358.

XX 01-MAY-1992; 92US-0877675.

XX (AMCY) AMERICAN CYANAMID CO.

PI Jacobsen JS, Vitek MP;

XX WPI; 1993-406194/51.

DR N-PSDB; AAQ54260.

XX New mutant forms of amyloid precursor protein - for detecting
 PT cpds. that modify activity of enzymes involved in precursor
 PT cleavage, also new nucleic acid encoding them

PS Disclosure; Page 34; 66pp; English.

XX Recombinant polypeptides produced using the coding sequences of
 CC mutant forms of amyloid precursor proteins comprising from the 5' to
 CC the 3' end a sequence encoding a marker and either (1) a sequence
 CC encoding the N-terminus of an amyloid precursor protein (APP) up to,
 CC but not including, the nucleotides encoding the beta amyloid protein
 CC (BAP) domain or (2) the BAP domain; or the two ligated together, can
 CC be used to detect drugs or compounds that inhibit/augment the
 CC activity of proteolytic enzymes which cleave APP to generate BAP
 CC fragments (deposition of which occurs in patients with Alzheimers
 CC disease and Down's syndrome). This fragment corresponding to amino
 CC acid residues 14-20 of BAP has been altered and APP's containing
 CC the altered BAP sequence show 0% secretion compared with those
 CC containing the wild type BAP sequence.

XX Sequence 7 AA;

Query Match 61.8%; Score 34; DB 14; Length 7;
 Best Local Similarity 85.7%; Pred. No. 6.4e+05;
 Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFF 8

Db 1 HOELVFF 7

RESULT 24

AAR87921

ID AAR87921 standard; peptide; 7 AA.

XX AAR87921;

XX 01-MAR-1996 (first entry)

XX Test peptide used in study of antagonism of amyloid beta protein.

DE amnesia; amyloid beta; Alzheimer's disease.

XX Synthetic.

XX WO9508999-Al.

PD 06-APR-1995.

XX

ID AAW02312 standard; peptide; 7 AA.
 XX AAW02312;
 AC
 DT 02-MAY-1997 (first entry)
 XX
 DE Beta-amyloid modulator peptide #3.
 XX
 XX Beta-amyloid; modulator; amyloid plaque; brain lesion; amyloidosis;
 KW cerebral blood vessel; Alzheimer's disease; amyloidogenic protein;
 KW familial amyloid polyneuropathy; familial amyloid cardiomyopathy;
 KW isolated cardiac amyloidosis; systemic senile amyloidosis; insulinoma;
 KW bovine spongiform encephalopathy; Creutzfeldt-Jakob disease; urticaria;
 KW adult-onset diabetes; familial Mediterranean fever; therapy; deafness;
 KW scrapie; familial amyloid nephropathy; hereditary cerebral haemorrhage.
 XX
 OS Synthetic.
 XX
 XX WO9628471-A1.
 PN
 XX
 XX 19-SEP-1996.
 PD
 XX
 XX 14-MAR-1996; 96WO-US03492.
 PF
 XX
 XX 27-OCT-1995; 95US-0548998.
 PR
 XX 14-MAR-1995; 95US-0404831.
 PR
 XX 07-JUN-1995; 95US-0475579.
 PR
 XX (PHAR-) PHARM PEPTIDES INC.
 PA
 XX Benjamin H, Chin J, Findeis MA, Garnick MB, Geffer ML;
 PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
 PI Molineaux S, Musso G, Reed MJ, Signer ER, Wakefield J;
 XX
 XX WPI; 1996-433762/43.
 DR
 XX
 XX Modulators of amyloid aggregation - comprising, e.g. amyloidogenic
 PT protein coupled (indirectly to at least 1 modifying gp., useful in
 PT treatment of Alzheimer's disease
 PT
 XX
 XX Claim 16; Page 91; 106pp; English.
 PS
 XX
 XX AAW02310-W02332 represent the peptide portions of the beta-amyloid
 CC modulator compounds of the invention. Beta-amyloid peptide is a 4
 CC kilodalton peptide that is the major protein component of amyloid
 CC plaques. Amyloid plaques are present both in the brain lesions, and in
 CC the walls of cerebral blood vessels in Alzheimer's disease patients.
 CC The amyloid modulators of the invention comprise an amyloidogenic protein
 CC or peptide (such as this sequence) coupled directly or indirectly to at
 CC least one modifying group. The modifying group is preferably a cyclic,
 CC heterocyclic, or polycyclic group, such as decalin, a cholanyl group, a
 CC biotin containing group, or a fluorescein containing group. These
 CC compounds then modulate the aggregation of these sequences to natural
 CC amyloid proteins or peptides when contacted with the natural
 CC amyloidogenic proteins or peptides. The modulator compounds can be used
 CC in the treatment of disorders associated with amyloidosis, such as
 CC familial amyloid polyneuropathy, familial amyloid cardiomyopathy,
 CC isolated cardiac amyloidosis, systemic senile amyloidosis, scrapie,
 CC bovine spongiform encephalopathy, Creutzfeldt-Jakob disease, adult-onset
 CC diabetes, insulinoma, familial Mediterranean fever, familial amyloid
 CC nephropathy with urticaria and deafness, hereditary cerebral haemorrhage
 CC and other types of amyloidosis. The modulators are also useful for the
 CC treatment of disorders associated with beta-amyloidosis, especially
 CC Alzheimer's disease.
 XX
 SQ Sequence 7 AA;
 Query Match 61.8%; Score 34; DB 17; Length 7;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 3 QKLVFFA 9
 | | | | | | |

Db 1 QKLVFFA 7
 RESULT 28
 AAW45942
 ID AAW45942 standard; peptide; 7 AA.
 XX
 AC AAW45942;
 AC
 XX 30-JUN-1998 (first entry)
 DT
 XX
 XX Amyloid beta peptide fragment.
 DE
 XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.
 KW
 XX Homo sapiens.
 OS
 XX WO9721728-A1.
 PN
 XX
 XX 19-JUN-1997.
 PD
 XX
 XX 09-DEC-1996; 96WO-SE01621.
 PF
 XX
 XX 29-DEC-1995; 95US-0009386.
 PR
 XX 12-DEC-1995; 95SE-0004467.
 PR
 XX (KARO-) KAROLINSKA INNOVATIONS AB.
 PA
 XX
 XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 PI
 XX WPI; 1997-332723/30.
 DR
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 PT
 XX
 XX Example 1; Figure 2B; 31pp; English.
 PS
 XX
 XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 7 AA;
 Query Match 61.8%; Score 34; DB 18; Length 7;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLV 6
 | | | | |
 Db 2 HHQKLV 7
 RESULT 29
 AAW49755
 ID AAW49755 standard; peptide; 7 AA.
 XX
 AC AAW49755;
 AC
 XX 12-OCT-1998 (first entry)
 DT
 XX

```

DE  Glutamine donor peptide.
XX
KW  Protein polymer; adhesive sealant; wound healing; cross-linking.
XX
XX  Synthetic.
OS
XX  US5773577-A.
PN
XX  30-JUN-1998.
PD
XX
XX  03-MAR-1994; 94US-0205518.
PF
XX
XX  02-MAR-1995; 95US-0397633.
PR
PR  03-MAR-1994; 94US-0205518.
XX
XX  (PROT-) PROTEIN POLYMER TECHNOLOGIES INC.
PA
XX  Cappello J;
PI
XX
XX  WPI; 1998-387091/33.
DR
XX
XX  New recombinant protein polymers - containing naturally occurring
PT  repetitive units for crosslinking by enzymes, useful as medical
PT  adhesives and sealants, depots and matrices
XX
XX  Example 9; Column 49; 70pp; English.
XX
XX  This is an example of a glutamine donor peptide that can be
CC  utilised in novel recombinant protein polymers of the invention.
CC  Such polymers (see AAW49710-28) typically comprise a repetitive
CC  amino acid backbone of repetitive units having a collagen, fibroin,
CC  elastin or keratin motif and at least 2 enzyme recognition
CC  sequences comprising a glutamine and/or lysine capable of enzyme
CC  catalysed isopeptide formation. The polymers are capable of
CC  covalent crosslinking by enzymatic reaction to form products which
CC  set quickly and have good adhesive properties and high strength.
CC  They can be used as medical adhesives and sealants, in the closure
CC  of wounds and repair of damaged tissues, prosthesis coatings, drug
CC  depots, and matrices for the transplantation of cells.
XX
XX  Sequence 7 AA;
SQ
Query Match 61.8%; Score 34; DB 19; Length 7;
Best Local Similarity 100.0%; Pred. NO. 6.4e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY  1 HHOKLV 6
Db  2 HHOKLV 7
|||||

RESULT 30
AAW89376
ID  AAW89376 standard; peptide; 7 AA.
XX
AC  AAW89376;
XX
XX  02-MAR-1999 (first entry)
DT
XX
XX  Beta-amyloid peptide derivative A-beta-15-21.
DE
XX
KW  Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
KW  aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
KW  familial amyloid polynuropathy; bovine spongiform encephalopathy;
KW  Creutzfeldt-Jakob disease; BAP.
XX
XX  Homo sapiens.
OS
XX  Synthetic.
XX
XX  US5854204-A.
PN
XX
XX  29-DEC-1998.
PD
XX

```

```

PF  14-MAR-1996; 96US-0612785.
XX
XX  14-MAR-1996; 96US-0612785.
PR  14-MAR-1995; 95US-0404831.
PR  07-JUN-1995; 95US-0475579.
PR  27-OCT-1995; 95US-0548998.
XX
XX  (PRAE-) PRAECIS PHARM INC.
PA
XX
XX  Benjamin H, Chin J, Findeis MA, Garnick MB, Geffer ML;
XX  Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
PI  Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;
XX
XX  WPI; 1999-094964/08.
DR
XX
XX  New peptide(s) derived from beta-amyloid peptide that inhibit
PT  amyloid aggregation - and neurotoxicity, specifically for treatment
PT  and prevention of Alzheimer's disease
XX
XX  Example 12; Column 64; 52pp; English.
XX
XX  The present invention describes beta-amyloid peptide (bAP) derivatives.
CC  The bAP derivatives inhibit aggregation of amyloidogenic proteins and
CC  peptides, specifically bAP, and their neurotoxicity, so are useful for
CC  treating and preventing any disease involving amyloidosis, specifically
CC  Alzheimer's disease but also Down's syndrome, familial amyloid
CC  polynuropathy or cardiomyopathy. The bAP derivatives are also used to diagnose
CC  Creutzfeldt-Jakob disease. The bAP derivatives are also used to diagnose
CC  these diseases, in vitro or in vivo, by detecting binding of bAP to
CC  labelled bAP derivatives. Some bAP derivatives inhibit bAP aggregation
CC  even when bAP is present in molar excess. The present sequence
CC  represents a bAP derivative.
XX
XX  Sequence 7 AA;
SQ
Query Match 61.8%; Score 34; DB 20; Length 7;
Best Local Similarity 100.0%; Pred. NO. 6.4e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY  3 QKLVFFA 9
Db  1 QKLVFFA 7
|||||

RESULT 31
AAB67281
ID  AAB67281 standard; peptide; 7 AA.
XX
XX  AAB67281;
AC
XX
XX  20-APR-2001 (first entry)
DT
XX
XX  Residues 16-22 of Alzheimer's Abeta peptide.
DE
XX
XX  Alzheimer's; Abeta; beta-strand.
KW
XX
XX  Homo sapiens.
OS
XX  WO200107473-A1.
PN
XX
XX  01-FEB-2001.
PD
XX
XX  28-JUL-2000; 2000WO-GB02901.
PF
XX
XX  28-JUL-1999; 99GB-0017724.
PR
XX
XX  (STOT/) STOTT K.
PA
XX  Stott K;
PI
XX
XX  WPI; 2001-182777/18.
DR
XX
XX  Novel chemical compound or composition useful for preventing
PT

```

PT beta-strand association, comprises peptides containing N-alpha
PT substituted L-amino acids

PS Claim 17; Page 46; 77pp; English.

XX The present invention relates to a chemical compound or composition
CC comprising a peptide with a beta strand forming section and
CC associates with a target beta-strand formed by a separate
CC peptide-containing molecule. The invention is useful for
CC inhibiting or reversing the association of target beta-strand,
CC formed by Alzheimer's Abeta peptide into a beta-sheet or beta-fibre
CC and the aggregation of proteins or peptides.

XX Sequence 7 AA;

Query Match 61.8%; Score 34; DB 22; Length 7;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVEFAE 10
|||||||

Db 1 KLVEFAE 7

RESULT 32

AAW45939
ID AAW45939 standard; peptide; 8 AA.

XX AAW45939;

DT 30-JUN-1998 (first entry)

XX Amyloid beta peptide fragment.

XX Anyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

XX WO9721728-A1.

XX 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

XX 29-DEC-1995; 95US-0009386.

XX 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

XX WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of
PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
PT Down's syndrome associated with amyloidosis.

PS Example 1; Figure 2B; 3lpp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The
CC invention relates to the use of peptide compounds for inhibition of
CC polymerisation of amyloid beta peptide (ABP), as model substances for
CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
CC tool for the identification of other organic compounds with similar
CC functional properties, or as ligands in positron emission tomography.
CC The peptides may be used in treatment of amyloidosis, especially in
CC treatment of Alzheimer's disease associated with amyloidosis, for
CC treatment or prevention of demens in patients with Down's syndrome, for
CC amyloidosis (Dutch type) or for the prevention of fibril formation of
CC human amyloid protein. They can also be used for identifying other
CC molecules with similar properties and/or as ligands for detection of

CC amyloid deposits using e.g. positron emission tomography.

SQ Sequence 8 AA;

Query Match 61.8%; Score 34; DB 18; Length 8;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
|||||||

Db 3 HHQKLV 8

RESULT 33

AAW32551

ID AAW32551 standard; peptide; 8 AA.

XX AAW32551;

DT 21-JAN-1998 (first entry)

XX Amyloidogenic sequence amyloid beta-peptide.

XX Anti-amyloid peptide; iAbeta; abnormal protein folding inhibitor;
KW Alzheimer's disease; dementia; Down's syndrome; amyloidosis disorder;
KW human prion disease; Kuru; Creutzfeldt-Jakob disease;
KW Gerstmann-Straussler-Scheinker Syndrome; animal prion disease;
KW prion associated human neurodegenerative disease; scrapie;
KW spongiform encephalopathy; transmissible mink encephalopathy;
KW chronic wasting disease; mule; deer; elk; human.

XX Homo sapiens.

OS Synthetic.

XX WO9639834-A1.

XX 19-DEC-1996.

XX 06-JUN-1996; 96WO-US10220.

XX 10-APR-1996; 96US-0630645.

XX 07-JUN-1995; 95US-0478326.

XX (UYNV) UNIV NEW YORK STATE.

XX Baumann MH, Frangione B, Soto-Jara C;

XX WPI; 1997-051637/05.

XX New inhibitors of fibrillogenesis proteins or peptides - used for
PT preventing, treating or detecting amyloidosis disorders such as
PT Alzheimer's disease.

PS Disclosure; Fig 1A; 63pp; English.

XX A method has been developed for the prevention or treatment of a
CC disorder or disease associated with the formation of amyloid or
CC amyloid-like deposits, involving the abnormal folding of a protein
CC or peptide. The method involves administering an inhibitory peptide
CC which prevents the abnormal folding or which dissolves existing amyloid
CC or amyloid-like deposits, where the peptide comprises a sequence of
CC 3-15 amino acid residues and has a hydrophobic cluster of at least 3
CC amino acids, where at least one of the 3 amino acids is a beta-sheet
CC blocking amino acid residue selected from Pro, Gly, Asn and His. The
CC present sequence represents an amyloidogenic sequence, amyloid beta-
CC peptide, which is involved in the formation of several amyloid deposits.
CC The inhibitory peptide is capable of associating with a structural
CC determinant on the protein or peptide to structurally block and inhibit
CC the abnormal folding into amyloid or amyloid-like deposits. The method
CC can be used for preventing, treating or detecting e.g. Alzheimer's
CC dementia or disease, Down's syndrome, other amyloidosis disorders,
CC human prion diseases such as Kuru, Creutzfeldt-Jakob disease, Gerstmann-
CC Straussler-Scheinker syndrome, prion associated human neurodegenerative

CC diseases or animal prion diseases such as scrapie, spongiform
 CC encephalopathy, transmissible mink encephalopathy and chronic wasting
 CC disease of mule deer and elk.

XX Sequence 8 AA;

Query Match 61.8%; Score 34; DB 18; Length 8;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
 |||||
 Db 1 KLVFFAE 7

RESULT 34

AAE10663
 ID AAE10663 standard; peptide; 8 AA.

XX AC AAE10663;

XX 10-DEC-2001 (first entry)

XX Human amyloid precursor protein substrate alpha-secretase peptide #2.

XX Human; aspartyl protease 1; Aspl; amyloid precursor protein; APP;
 KW Alzheimer's disease; AD; dementia; neurofibrillary tangle; gliosis;
 KW amyloid plaque; neuronal loss; proteolytic; neurotropic; neuroprotective;
 alpha-secretase.

XX Homo sapiens.

XX Key Location/Qualifiers
 FT Cleavage-site 4..5

PN GB2357767-A.

XX 04-JUL-2001.

XX 22-SEP-2000; 2000GB-0023315.

XX 23-SEP-1999; 99US-0155493.

XX 23-SEP-1999; 99US-0404133.

XX 23-SEP-1999; 99WO-US20881.

XX 13-OCT-1999; 99US-0416901.

XX 06-DEC-1999; 99US-0169232.

XX (PHAA) PHARMACIA & UPJOHN CO.

XX Bienkowski MJ, Gurney M;

XX WPI; 2001-444208/48.

XX Polypeptide comprising fragments of human aspartyl protease with
 PT amyloid precursor protein processing activity and alpha-secretase
 PT activity, for identifying modulators useful in treating Alzheimer's
 PT disease -

XX Claim 10; Page 163; 187pp; English.

XX The patent discloses human aspartyl protease 1 (hu-Aspl) or modified
 CC Aspl proteins which lack transmembrane domain or amino terminal
 CC domain or cytoplasmic domain and retains alpha-secretase activity
 CC and amyloid protein precursor (APP) processing activity. The proteins
 CC of the invention are useful for assaying hu-Aspl alpha-secretase
 CC activity, which in turn is useful for identifying modulators of
 CC hu-Aspl alpha-secretase activity, where modulators that increase
 CC hu-Aspl alpha-secretase activity are useful for treating Alzheimer's
 CC disease (AD) which causes progressive dementia with consequent
 CC formation of amyloid plaques, neurofibrillary tangles, gliosis and
 CC neuronal loss. Hu-Aspl protease substrate is useful for assaying
 CC hu-Aspl proteolytic activity, by contacting hu-Aspl protein with
 CC the substrate under acidic conditions and determining the level of

CC hu-Aspl proteolytic activity. The present sequence is human amyloid
 CC precursor protein (APP) substrate alpha-secretase peptide which is
 CC used for determining the enzymatic activity of Asp-1 protein lacking
 CC transmembrane domain (TM) and containing a (His)6 tag.

XX Sequence 8 AA;

Query Match 61.8%; Score 34; DB 22; Length 8;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
 |||||
 Db 1 KLVFFAE 7

RESULT 35

AAE02615
 ID AAE02615 standard; peptide; 8 AA.

XX AC AAE02615;

XX 10-AUG-2001 (first entry)

XX Human amyloid precursor protein substrate alpha-secretase peptide #2.

XX Human; alpha-secretase; amyloid precursor protein; APP; therapy;
 KW Alzheimer's disease; antialzheimer's; aspartyl protease 1; Aspl;
 KW beta-secretase.

XX Homo sapiens.

XX Key Location/Qualifiers
 FT Cleavage-site 4..5

PN WO200123533-A2.

XX 05-APR-2001.

XX 22-SEP-2000; 2000WO-US26080.

XX 23-SEP-1999; 99US-0155493.

XX 23-SEP-1999; 99WO-US20881.

XX 13-OCT-1999; 99US-0416901.

XX 06-DEC-1999; 99US-0169232.

XX (PHAA) PHARMACIA & UPJOHN CO.

XX Gurney M, Bienkowski MJ;

XX WPI; 2001-290516/30.

XX Enzymes that cleave the alpha-secretase site of the amyloid precursor
 PT protein, useful for the treatment of Alzheimer's disease -

XX Claim 10; Page 98; 189pp; English.

XX The present invention relates to enzymes for cleaving the alpha-
 CC secretase site of the amyloid precursor protein (APP) and methods of
 CC identifying those enzymes. The methods may be used to identify enzymes
 CC that may be used to cleave the alpha-secretase cleavage site of the APP
 CC protein. The enzymes may be used to treat or modulate the progress of
 CC Alzheimer's disease. The present sequence is human amyloid precursor
 CC protein (APP) substrate alpha-secretase peptide which is used for
 CC determining the enzymatic activity of Asp-1 deltatm (His)6 protein.

XX Sequence 8 AA;

Query Match 61.8%; Score 34; DB 22; Length 8;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10

```

Db      1 KLVFFAE 7
      |||||
RESULT 36
AAY79938
ID AAY79938 standard; peptide; 10 AA.
XX
AC AAY79938;
XX
DT 11-MAY-2000 (first entry)
XX
DE Beta-amyloid recognition peptide SEQ ID NO:3.
XX
KW Beta-amyloid; inhibitor; recognition element; hybrid; aggregation;
KW Alzheimer's disease; neuroprotective; nootropic.
XX
OS Homo sapiens.
XX
PN US6022859-A.
XX
PD 08-FEB-2000.
XX
PF 14-NOV-1997; 97US-0970833.
XX
PR 15-NOV-1996; . 96US-0030840.
XX
PA (WISC ) WISCONSIN ALUMNI RES FOUND.
XX
PI Murphy RM, Kiessling LL;
XX
DR WPI; 2000-160387/14.
XX
PT Beta-amyloid inhibitor useful for treating Alzheimer's disease -
XX
PS Example; Column 7; 15pp; English.
XX
CC The present invention describes a beta-amyloid inhibitor peptide.
CC Beta-amyloid inhibitors have neuroprotective and nootropic
CC properties. The inhibitor peptides are useful for the treatment of
CC Alzheimer's disease. The present sequence represents a beta-amyloid
CC recognition peptide used in the exemplification of present invention.
XX
SQ Sequence 10 AA;
      Query Match 61.8%; Score 34; DB 21; Length 10;
      Best Local Similarity 100.0%; Pred. No. 2;
      Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy      4 KLVFFAE 10
      |||||
Db      1 KLVFFAE 7

RESULT 37
AAB46221
ID AAB46221 standard; peptide; 10 AA.
XX
AC AAB46221;
XX
DT 04-APR-2001 (first entry)
XX
DE Human APP derived immunogenic peptide #17.
XX
KW Amyloid deposit; APP; Abeta; brain; human; clearing response; nootropic;
KW Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
KW amyloid precursor protein; Alzheimer's disease.
XX
OS Homo sapiens.
XX
PN WO200072880-A2.
XX
PD 07-DEC-2000.

XX
XX 26-MAY-2000; 2000WO-US14810.
XX
XX 28-MAY-1999; 99US-0322289.
XX
XX (NEUR-) NEURALAB LTD.
XX
XX Schenk DB, Bard F, Vasquez NJ, Yednock T;
XX
XX WPI; 2001-032104/04.
XX
XX Preventing or treating a disease associated with amyloid deposits,
XX especially Alzheimer's disease, comprises administering amyloid
XX specific antibody -
XX
XX Disclosure; Figure 19; 143pp; English.
XX
XX This invention describes a novel method of preventing or treating a
XX disease associated with amyloid deposits of amyloid precursor protein
XX (APP) Abeta fragments in the brain of a patient, which comprises
XX administering to the patient: (a) an antibody that binds to Abeta, the
XX antibody binds to an amyloid deposit and induces a clearing response (Fc
XX receptor mediated phagocytosis) against it (b) a polypeptide containing
XX an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
XX Abeta. The products of the invention have nootropic and neuroprotective
XX activity. The method is also useful for monitoring a course of treatment
XX being administered to a patient e.g. active and passive immunization. The
XX methods are useful for prophylactic and therapeutic treatment of
XX Alzheimer's disease.
XX
XX Sequence 10 AA;
      Query Match 61.8%; Score 34; DB 22; Length 10;
      Best Local Similarity 100.0%; Pred. No. 2;
      Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy      1 HHQKLV 6
      |||||
Db      5 HHQKLV 10

RESULT 38
AAB46228
ID AAB46228 standard; peptide; 10 AA.
XX
AC AAB46228;
XX
XX 04-APR-2001 (first entry)
XX
XX Human APP derived immunogenic peptide #24.
XX
XX Amyloid deposit; APP; Abeta; brain; human; clearing response; nootropic;
XX Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
XX amyloid precursor protein; Alzheimer's disease.
XX
XX Homo sapiens.
XX
XX WO200072880-A2.
XX
XX 07-DEC-2000.
XX
XX 26-MAY-2000; 2000WO-US14810.
XX
XX 28-MAY-1999; 99US-0322289.
XX
XX (NEUR-) NEURALAB LTD.
XX
XX Schenk DB, Bard F, Vasquez NJ, Yednock T;
XX
XX WPI; 2001-032104/04.
XX
XX Preventing or treating a disease associated with amyloid deposits,
XX

```

PT especially Alzheimer's disease, comprises administering amyloid
PT specific antibody -
PS Disclosure; Figure 19; 143pp; English.
XX
XX This invention describes a novel method of preventing or treating a
CC disease associated with amyloid deposits of amyloid precursor protein
CC (APP) Abeta fragments in the brain of a patient, which comprises
CC administering to the patient: (a) an antibody that binds to Abeta, the
CC antibody binds to an amyloid deposit and induces a clearing response (Fc
CC receptor mediated phagocytosis) against it (b) a polypeptide containing
CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
CC that induces an immunogenic response against residues 1-3 to 7-11 of
CC Abeta. The products of the invention have neurotropic and neuroprotective
CC activity. The method is also useful for monitoring a course of treatment
CC being administered to a patient e.g. active and passive immunization. The
CC methods are useful for prophylactic and therapeutic treatment of
CC Alzheimer's disease.
XX
XX Sequence 10 AA;
SQ
Query Match 61.8%; Score 34; DB 22; Length 10;
Best Local Similarity 100.0%; Pred. No. 2;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 4 KLVFFAE 10
DB 1 KLVFFAE 7
|||||
RESULT 39
AAW45945
ID AAW45945 standard; peptide; 6 AA.
XX
XX AAW45945;
AC
DT 30-JUN-1998 (first entry)
XX
XX Amyloid beta peptide fragment.
DE
XX
XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
KW positron emission tomography; PET; Down's syndrome; amyloidosis.
KW
OS Homo sapiens.
OS
XX
XX WO9721728-A1.
PN
XX
XX 19-JUN-1997.
PD
XX
XX 09-DEC-1996; 96WO-SE01621.
PF
XX
XX 29-DEC-1995; 95US-0009386.
PR
XX
XX 12-DEC-1995; 95SE-0004467.
PR
XX
XX (KARO-) KAROLINSKA INNOVATIONS AB.
PA
XX
XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
PI WPI; 1997-332723/30.
XX
XX Use of new and known peptide(s) for inhibition of polymerisation of
PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
PT Down's syndrome associated with amyloidosis.
PT
XX
XX Example 1; Figure 2B; 31pp; English.
PS
XX
XX This sequence represents a fragment of the amyloid beta peptide. The
CC invention relates to the use of peptide compounds for inhibition of
CC polymerisation of amyloid beta peptide (ABP), as model substances for
CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
CC tool for the identification of other organic compounds with similar
CC functional properties, or as ligands in positron emission tomography.
CC The peptides may be used in treatment of amyloidosis, especially in

CC treatment of Alzheimer's disease associated with amyloidosis, for
CC treatment or prevention of demens in patients with Down's syndrome, for
CC treatment or prevention of hereditary cerebral haemorrhage with
CC amyloidosis (Dutch type) or for the prevention of fibril formation of
CC human amyloid protein. They can also be used for identifying other
CC molecules with similar properties and/or as ligands for detection of
CC amyloid deposits using e.g. positron emission tomography.
XX
SQ Sequence 6 AA;
Query Match 58.2%; Score 32; DB 18; Length 6;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2 HOKLVF 7
DB 1 HOKLVF 6
|||||
RESULT 40
AAR45233
ID AAR45233 standard; Peptide; 7 AA.
XX
XX AAR45233;
AC
XX 20-JUN-1994 (first entry)
DT
XX
XX Beta amyloid protein fragment.
DE
XX
XX Amyloid precursor protein; APP; beta amyloid protein; BAP;
KW detection; Alzheimer's disease; Down's syndrome.
KW
XX
XX Homo sapiens.
OS
XX
XX AU9338358-A.
PN
XX
XX 04-NOV-1993.
PD
XX
XX 03-MAY-1993; 93AU-0038358.
PF
XX
XX 01-MAY-1992; 92US-0877675.
PR
XX
XX (AMCY) AMERICAN CYANAMID CO.
PA
XX
XX Jacobsen JS, Vitek MP;
PI
XX
XX WPI; 1993-406194/51.
DR
XX
XX N-PSDB; AAQ54261.
DR
XX
XX New mutant forms of amyloid precursor protein - for detecting
PT cpds. that modify activity of enzymes involved in precursor
PT cleavage, also new nucleic acid encoding them
PT
XX
XX Disclosure; Page 34; 66pp; English.
PS
XX
XX Recombinant polypeptides produced using the coding sequences of
CC mutant forms of amyloid precursor proteins comprising from the 5' to
CC the 3' end a sequence encoding a marker and either (1) a sequence
CC encoding the N-terminus of an amyloid precursor protein (APP) up to,
CC but not including, the nucleotides encoding the beta amyloid protein
CC (BAP) domain or (2) the BAP domain; or the two ligated together, can
CC be used to detect drugs or compounds that inhibit/augment the
CC activity of proteolytic enzymes which cleave APP to generate BAP
CC fragments (deposition of which occurs in patients with Alzheimer's
CC disease and Down's syndrome). This fragment corresponding to amino
CC acid residues 14-20 of BAP has been altered and APP's containing
CC the altered BAP sequence show 10-20% secretion compared with those
CC containing the wild type BAP sequence.
XX
XX Sequence 7 AA;
Query Match 56.4%; Score 31; DB 14; Length 7;
Best Local Similarity 85.7%; Pred. No. 6.4e+05;

Matches 6; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HOKLVFF 8
 Db 1 HQVLVFF 7

RESULT 41
 AAB82639
 ID AAB82639 standard; Peptide; 7 AA.
 XX
 AC AAB82639;
 XX
 DT 02-OCT-2001 (first entry)
 XX
 DE All-D peptide used in Alzheimer's disease vaccine.
 XX
 KW Alzheimer's disease; amyloidosis; amyloid-related disease;
 KW vaccine; therapy; antigen.
 XX
 OS Synthetic.
 XX
 FH Key Location/Qualifiers
 FT Misc-difference 1..7 /note= "all D-form residues"
 XX
 PN WO200139796-A2.
 XX
 PD 07-JUN-2001.
 XX
 PF 29-NOV-2000; 2000WO-CA01413.
 XX
 PR 29-NOV-1999; 99US-0168594.
 PR 28-NOV-2000; 2000US-0724842.
 XX
 PA (NEUR-) NEUROCHEM INC.
 XX
 PI Chalfour R, Hebert L, Kong X, Gervais F;
 XX
 DR WPI; 2001-441458/47.
 XX
 PT Preventing/treating amyloid-related disease, especially Alzheimer's
 PT disease, comprises administering antigenic all-D peptide, e.g. as
 PT vaccine, which elicits production of antibodies to prevent
 PT fibrillogenesis and associated cellular toxicity -
 XX
 PS Disclosure; Page 11; 31pp; English.
 CC
 CC The present sequence is that of an all-D peptide suitable for
 CC use in preparing vaccines for preventing or treating Alzheimer's
 CC disease and other amyloid related disorders in humans. It is based
 CC on a portion of amyloid-beta peptide (see AAB82622), and may be
 CC modified by removing or inserting 1 or more amino acid residues, or
 CC by substituting 1 or more amino acid residues with other amino acid
 CC residues or non-amino acid fragments. Vaccines of the invention
 CC are produced using 'non-self' peptides synthesised from the
 CC unnatural D-configuration amino acids to avoid the drawbacks of
 CC 'self' proteins. The all-D peptides need not be aggregated to be
 CC operative or immunogenic. They preferably interact with at
 CC least 1 region of an amyloid protein, e.g. the beta-sheet region
 CC or GAG-binding site region, the amyloid-beta peptide, or their
 CC immunogenic fragments, protein conjugates, immunogenic derivative
 CC peptides and immunogenic peptidomimetics. Examples include all-D
 CC peptides corresponding to residues 1-42, 1-40, 1-35, 1-28, 1-7,
 CC 10-16, 16-21 and 36-42 of the amyloid-beta peptide and the all-D
 CC derivative peptides given in AAB82623-64. The vaccine elicits a
 CC preferential TH-2 or TH-1 response, preventing fibrillogenesis and
 CC associated cellular toxicity. The amyloid related diseases may be
 CC localised amyloidosis, e.g. diabetes type II, neurodegenerative
 CC diseases, e.g. bovine spongiform encephalitis, Creutzfeldt-Jakob
 CC disease, scrapie, cerebral amyloid angiopathy, and prion protein
 CC related disorders, or systemic amyloidosis associated with chronic
 CC infection (e.g. tuberculosis) or chronic inflammation (e.g.

CC rheumatoid arthritis), familial Mediterranean fever (FMF) and
 CC systemic amyloidosis found in long-term haemodialysis patients.

XX
 SQ Sequence 7 AA;
 Query Match 56.4%; Score 31; DB 22; Length 7;
 Best Local Similarity 85.7%; Pred. No. 6.4e+05;
 Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFFAE 10
 Db 1 KLVFFAQ 7

RESULT 42
 AAB82640
 ID AAB82640 standard; Peptide; 7 AA.
 XX
 AC AAB82640;
 XX
 DT 02-OCT-2001 (first entry)
 XX
 DE All-D peptide used in Alzheimer's disease vaccine.
 XX
 KW Alzheimer's disease; amyloidosis; amyloid-related disease;
 KW vaccine; therapy; antigen.
 XX
 OS Synthetic.
 XX
 FH Key Location/Qualifiers
 FT Misc-difference 1..7 /note= "all D-form residues"
 FT Modified-site 6 /note= "C-terminal amide"
 FT
 XX
 PN WO200139796-A2.
 XX
 PD 07-JUN-2001.
 XX
 PF 29-NOV-2000; 2000WO-CA01413.
 XX
 PR 29-NOV-1999; 99US-0168594.
 PR 28-NOV-2000; 2000US-0724842.
 XX
 PA (NEUR-) NEUROCHEM INC.
 XX
 PI Chalfour R, Hebert L, Kong X, Gervais F;
 XX
 DR WPI; 2001-441458/47.
 XX
 PT Preventing/treating amyloid-related disease, especially Alzheimer's
 PT disease, comprises administering antigenic all-D peptide, e.g. as
 PT vaccine, which elicits production of antibodies to prevent
 PT fibrillogenesis and associated cellular toxicity -
 XX
 PS Disclosure; Page 11; 31pp; English.
 CC
 CC The present sequence is that of an all-D peptide suitable for
 CC use in preparing vaccines for preventing or treating Alzheimer's
 CC disease and other amyloid related disorders in humans. It is based
 CC on a portion of amyloid-beta peptide (see AAB82622), and may be
 CC modified by removing or inserting 1 or more amino acid residues, or
 CC by substituting 1 or more amino acid residues with other amino acid
 CC residues or non-amino acid fragments. Vaccines of the invention
 CC are produced using 'non-self' peptides synthesised from the
 CC unnatural D-configuration amino acids to avoid the drawbacks of
 CC 'self' proteins. The all-D peptides need not be aggregated to be
 CC operative or immunogenic. They preferably interact with at
 CC least 1 region of an amyloid protein, e.g. the beta-sheet region
 CC or GAG-binding site region, the amyloid-beta peptide, or their
 CC immunogenic fragments, protein conjugates, immunogenic derivative
 CC peptides and immunogenic peptidomimetics. Examples include all-D
 CC peptides corresponding to residues 1-42, 1-40, 1-35, 1-28, 1-7,
 CC 10-16, 16-21 and 36-42 of the amyloid-beta peptide and the all-D
 CC derivative peptides given in AAB82623-64. The vaccine elicits a
 CC preferential TH-2 or TH-1 response, preventing fibrillogenesis and
 CC associated cellular toxicity. The amyloid related diseases may be
 CC localised amyloidosis, e.g. diabetes type II, neurodegenerative
 CC diseases, e.g. bovine spongiform encephalitis, Creutzfeldt-Jakob
 CC disease, scrapie, cerebral amyloid angiopathy, and prion protein
 CC related disorders, or systemic amyloidosis associated with chronic
 CC infection (e.g. tuberculosis) or chronic inflammation (e.g.

CC 10-16, 16-21 and 36-42 of the amyloid-beta peptide and the all-D
 CC derivative peptides given in AAB82623-64. The vaccine elicits a
 CC preferential TH-2 or TH-1 response, preventing fibrillogenesis and
 CC associated cellular toxicity. The amyloid related diseases may be
 CC localised amyloidosis, e.g. diabetes type II, neurodegenerative
 CC diseases, e.g. bovine spongiform encephalitis, Creutzfeldt-Jakob
 CC disease, scrapie, cerebral amyloid angiopathy, and prion protein
 CC related disorders, or systemic amyloidosis associated with chronic
 CC infection (e.g. tuberculosis) or chronic inflammation (e.g.
 CC rheumatoid arthritis), familial Mediterranean fever (FMF) and
 CC systemic amyloidosis found in long-term haemodialysis patients.
 XX
 XX
 SQ Sequence 7 AA;

Query Match 56.4%; Score 31; DB 22; Length 7;
 Best Local Similarity 85.7%; Pred. No. 6.4e+05;
 Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
 Db 1 KLVFFAQ 7
 |||||

RESULT 43
 AAB48491
 ID AAB48491 standard; Peptide; 7 AA.
 XX
 AC AAB48491;
 XX
 DT 02-MAR-2001 (first entry)
 XX
 DE Antifibrillogenic peptide #18.
 XX
 KW Nootropic; neuroprotective; antifibrillogenic; amyloidosis inhibition;
 KW cytoprotection; amyloid deposit degradation; amyloidosis disorder;
 KW Alzheimer's disease.
 XX
 OS Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Modified-site 7 /note- "C-terminal amide"
 FT
 XX
 PN WO200068263-A2.
 XX
 PD 16-NOV-2000.
 XX
 PF 04-MAY-2000; 2000WO-CA00515.
 XX
 PR 05-MAY-1999; 99US-0132592.
 XX
 PA (NEUR-) NEUROCHEM INC.
 XX
 PI Chalifour R, Gervais F, Gupta A;
 XX
 XX WPI; 2001-031852/04.
 XX
 DR Antifibrillogenic agent useful for inhibiting amyloidosis and/or for
 XX cytoprotection for treating amyloidosis disorders, comprises a peptide,
 XX its isomer or peptidomimetic -
 XX
 PT
 PT
 PT
 XX
 PS Claim 7; Page 25; 46pp; English.
 XX
 CC Peptides AAB48474-B48496 are antifibrillogenic agents that can be used
 CC for inhibiting amyloidosis and/or for cytoprotection. The peptides of
 CC AAB48474-B48496 cause the breakdown of amyloid deposits and are
 CC therefore useful for treating amyloidosis disorders such as Alzheimer's
 CC disease. Peptides AAB48474-B48496 were identified from the
 CC glycosaminoglycan binding region and the prot-prot interaction region of
 CC the human amyloid protein.
 XX
 XX
 SQ Sequence 7 AA;

Query Match 56.4%; Score 31; DB 22; Length 7;
 Best Local Similarity 85.7%; Pred. No. 6.4e+05;
 Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
 Db 1 KLVFFAQ 7
 |||||

RESULT 44
 AAB48492
 ID AAB48492 standard; Peptide; 7 AA.
 XX
 AC AAB48492;
 XX
 DT 02-MAR-2001 (first entry)
 XX
 DE Antifibrillogenic peptide #19.
 XX
 KW Nootropic; neuroprotective; antifibrillogenic; amyloidosis inhibition;
 KW cytoprotection; amyloid deposit degradation; amyloidosis disorder;
 KW Alzheimer's disease.
 XX
 OS Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Modified-site 7 /note- "C-terminal amide"
 FT
 XX
 PN WO200068263-A2.
 XX
 PD 16-NOV-2000.
 XX
 PF 04-MAY-2000; 2000WO-CA00515.
 XX
 PR 05-MAY-1999; 99US-0132592.
 XX
 PA (NEUR-) NEUROCHEM INC.
 XX
 PI Chalifour R, Gervais F, Gupta A;
 XX
 XX WPI; 2001-031852/04.
 XX
 DR Antifibrillogenic agent useful for inhibiting amyloidosis and/or for
 XX cytoprotection for treating amyloidosis disorders, comprises a peptide,
 XX its isomer or peptidomimetic -
 XX
 PT
 PT
 PT
 XX
 PS Claim 7; Page 25; 46pp; English.
 XX
 CC Peptides AAB48474-B48496 are antifibrillogenic agents that can be used
 CC for inhibiting amyloidosis and/or for cytoprotection. The peptides of
 CC AAB48474-B48496 cause the breakdown of amyloid deposits and are
 CC therefore useful for treating amyloidosis disorders such as Alzheimer's
 CC disease. Peptides AAB48474-B48496 were identified from the
 CC glycosaminoglycan binding region and the prot-prot interaction region of
 CC the human amyloid protein.
 XX
 XX
 SQ Sequence 7 AA;

Query Match 56.4%; Score 31; DB 22; Length 7;
 Best Local Similarity 85.7%; Pred. No. 6.4e+05;
 Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
 Db 1 KLVFFAQ 7
 |||||

RESULT 45
 AAB45952
 ID AAB45952 standard; peptide; 5 AA.
 XX
 AC AAB45952;
 XX
 DT 30-JUN-1998 (first entry)
 XX
 DE Amyloid beta peptide fragment.

XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.
 XX Homo sapiens.

XX WO9721728-A1.

XX 19-JUN-1997.

XX 09-DEC-1996; 96WO-SE01621.

XX 29-DEC-1995; 95US-0009386.

XX 12-DEC-1995; 95SE-0004467.

XX (KARO-) KAROLINSKA INNOVATIONS AB.

XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;

XX WPI; 1997-332723/30.

XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.

XX Example 1; Figure 2B; 31pp; English.

XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of dementia in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.

XX Sequence 5 AA;

Query Match 54.5%; Score 30; DB 18; Length 5;

Best Local Similarity 100.0%; Pred. No. 6.4e+05;

Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKL 5

DB 1 HHQKL 5

RESULT 46

AAW02313
 ID AAW02313 standard; peptide; 6 AA.

XX AAW02313;

XX 02-MAY-1997 (first entry)

XX Beta-amyloid modulator peptide #4.

XX Beta-amyloid; modulator; amyloid plaque; brain lesion; amyloidosis;
 KW cerebral blood vessel; Alzheimer's disease; amyloidogenic protein;
 KW familial amyloid polynuropathy; familial amyloid cardiomyopathy;
 KW isolated cardiac amyloidosis; systemic senile amyloidosis; insulinoma;
 KW bovine spongiform encephalopathy; Creutzfeldt-Jakob disease; urticaria;
 KW adult-onset diabetes; familial Mediterranean fever; therapy; deafness;
 KW scrapie; familial amyloid nephropathy; hereditary cerebral haemorrhage.

XX Synthetic.

XX

PN WO9628471-A1.

XX 19-SEP-1996.

XX 14-MAR-1996; 96WO-US03492.

XX 27-OCT-1995; 95US-0548998.

XX 14-MAR-1995; 95US-0404831.

XX 07-JUN-1995; 95US-0475579.

XX (PHAR-) PHARM PEPTIDES INC.

XX Benjamin H, Chin J, Findeis MA, Garnick MB, Gefter ML;

PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;

PI Mollineaux S, Musso G, Reed MJ, Signer ER, Wakefield J;

XX WPI; 1996-433762/43.

XX Modulators of amyloid aggregation - comprising, e.g. amyloidogenic
 PT protein coupled (in)directly to at least 1 modifying gp., useful in
 PT treatment of Alzheimer's disease

XX Claim 16; Page 91; 106pp; English.

XX AAW02310-W02332 represent the peptide portions of the beta-amyloid
 CC modulator compounds of the invention. Beta-amyloid peptide is a 4
 CC kilodalton peptide that is the major protein component of amyloid
 CC plaques. Amyloid plaques are present both in the brain lesions, and in
 CC the walls of cerebral blood vessels in Alzheimer's disease patients.
 CC The amyloid modulators of the invention comprise an amyloidogenic protein
 CC or peptide (such as this sequence) coupled directly or indirectly to at
 CC least one modifying group. The modifying group is preferably a cyclic,
 CC heterocyclic, or polycyclic group, such as decalin, a cholanyl group, a
 CC biotin containing group, or a fluorescein containing group. These
 CC compounds then modulate the aggregation of these sequences to natural
 CC amyloidogenic proteins or peptides when contacted with the natural
 CC amyloidogenic proteins or peptides. The modulator compounds can be used
 CC in the treatment of disorders associated with amyloidosis, such as
 CC familial amyloid polynuropathy, familial amyloid cardiomyopathy,
 CC isolated cardiac amyloidosis, systemic senile amyloidosis, scrapie,
 CC bovine spongiform encephalopathy, Creutzfeldt-Jakob disease, adult-onset
 CC diabetes, insulinoma, familial Mediterranean fever, familial amyloid
 CC nephropathy with urticaria and deafness, hereditary cerebral haemorrhage
 CC and other types of amyloidosis. The modulators are also useful for the
 CC treatment of disorders associated with beta-amyloidosis, especially
 CC Alzheimer's disease.

XX Sequence 6 AA;

Query Match 54.5%; Score 30; DB 17; Length 6;

Best Local Similarity 100.0%; Pred. No. 6.4e+05;

Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFF 8

DB 1 QKLVFF 6

RESULT 47

AAW45947
 ID AAW45947 standard; peptide; 6 AA.

XX AAW45947;

XX 30-JUN-1998 (first entry)

XX Amyloid beta peptide fragment.

XX Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.

XX Homo sapiens.

XX

PN WO9721728-A1.
 PD 19-JUN-1997.
 XX
 XX 09-DEC-1996; 96WO-SE01621.
 XX
 XX 29-DEC-1995; 95US-0009386.
 PR 12-DEC-1995; 95SE-0004467.
 XX
 XX (KARO-) KAROLINSKA INNOVATIONS AB.
 PA
 XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 XX WPI; 1997-332723/30.
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 XX
 XX Example 1; Figure 2B; 3lpp; English.
 PS
 XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 6 AA;
 Query Match 54.5%; Score 30; DB 18; Length 6;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKL 5
 Db |
 |
 2 HHQKL 6
 RESULT 48
 AAW45944
 ID AAW45944 standard; peptide; 6 AA.
 XX
 AC AAW45944;
 XX
 XX 30-JUN-1998 (first entry)
 DT
 XX
 DE Amyloid beta peptide fragment.
 XX
 KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.
 KW
 XX Homo sapiens.
 OS
 XX WO9721728-A1.
 PN
 XX 19-JUN-1997.
 PD
 XX 09-DEC-1996; 96WO-SE01621.
 PF
 XX 29-DEC-1995; 95US-0009386.
 PR
 XX 12-DEC-1995; 95SE-0004467.
 XX
 XX (KARO-) KAROLINSKA INNOVATIONS AB.
 PA
 XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 XX WPI; 1997-332723/30.
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 XX
 XX Example 1; Figure 2B; 3lpp; English.
 PS
 XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 6 AA;
 Query Match 54.5%; Score 30; DB 18; Length 6;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKL 5
 Db |
 |
 2 HHQKL 6
 RESULT 48
 AAW45944
 ID AAW45944 standard; peptide; 6 AA.
 XX
 AC AAW45944;
 XX
 XX 30-JUN-1998 (first entry)
 DT
 XX
 DE Amyloid beta peptide fragment.
 XX
 KW Amyloid beta peptide; Alzheimer's disease; polymerisation; aggregation;
 KW positron emission tomography; PET; Down's syndrome; amyloidosis.
 KW
 XX Homo sapiens.
 OS
 XX WO9721728-A1.
 PN
 XX 19-JUN-1997.
 PD
 XX 09-DEC-1996; 96WO-SE01621.
 PF
 XX 29-DEC-1995; 95US-0009386.
 PR
 XX 12-DEC-1995; 95SE-0004467.
 XX
 XX (KARO-) KAROLINSKA INNOVATIONS AB.
 PA
 XX Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 XX WPI; 1997-332723/30.
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 XX
 XX Example 1; Figure 2B; 3lpp; English.
 PS
 XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 6 AA;
 Query Match 54.5%; Score 30; DB 18; Length 6;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 3 QKLVFF 8
 Db |
 |
 1 QKLVFF 6
 RESULT 49
 AAY39801
 ID AAY39801 standard; peptide; 6 AA.
 XX
 AC AAY39801;
 XX
 XX 29-NOV-1999 (first entry)
 DT
 XX
 DE Beta-amyloid protein fragment.
 XX
 KW Beta-amyloid protein; Alzheimer's disease; amyloidosis; joint swelling;
 KW long-standing inflammation; malignancy; Familial Mediterranean Fever;
 KW multiple myeloma; plasma cell dyscrasia; long-term haemodialysis; Kuru;
 KW carpal tunnel syndrome; multiple spontaneous fracture; radiolucency;
 KW endocrine tumour; medullary carcinoma; Down's syndrome; scrapie;
 KW Creutzfeldt-Jakob disease; Gerstmann Strausler Syndrome;
 KW subacute spongiform encephalopathy; therapy.
 XX
 XX Homo sapiens.
 OS
 XX US5958883-A.
 PN
 XX 28-SEP-1999.
 PD
 XX 05-JUN-1995; 95US-0461216.
 PF
 XX 23-OCT-1992; 92US-0969734.
 PR
 XX 23-SEP-1992; 92US-0950417.
 XX
 XX (UNIW) UNIV WASHINGTON.
 PA
 XX Snow AD;
 PI
 XX WPI; 1999-561062/47.
 DR
 XX Peptides of 6-8 amino acids useful for treating or preventing
 PT amyloidosis -

PI Naslund J, Nordstedt C, Terenius L, Thyberg J, Tjernberg LO;
 XX WPI; 1997-332723/30.
 XX
 XX Use of new and known peptide(s) for inhibition of polymerisation of
 PT amyloid beta peptide - e.g. for treatment of Alzheimer's disease or
 PT Down's syndrome associated with amyloidosis.
 XX
 XX Example 1; Figure 2B; 3lpp; English.
 PS
 XX This sequence represents a fragment of the amyloid beta peptide. The
 CC invention relates to the use of peptide compounds for inhibition of
 CC polymerisation of amyloid beta peptide (ABP), as model substances for
 CC synthesis of ABP-ligands for inhibition of polymerisation of ABP, as a
 CC tool for the identification of other organic compounds with similar
 CC functional properties, or as ligands in positron emission tomography.
 CC The peptides may be used in treatment of amyloidosis, especially in
 CC treatment of Alzheimer's disease associated with amyloidosis, for
 CC treatment or prevention of demens in patients with Down's syndrome, for
 CC treatment or prevention of hereditary cerebral haemorrhage with
 CC amyloidosis (Dutch type) or for the prevention of fibril formation of
 CC human amyloid protein. They can also be used for identifying other
 CC molecules with similar properties and/or as ligands for detection of
 CC amyloid deposits using e.g. positron emission tomography.
 XX
 SQ Sequence 6 AA;
 Query Match 54.5%; Score 30; DB 18; Length 6;
 Best Local Similarity 100.0%; Pred. No. 6.4e+05;
 Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 3 QKLVFF 8
 Db |
 |
 1 QKLVFF 6
 RESULT 49
 AAY39801
 ID AAY39801 standard; peptide; 6 AA.
 XX
 AC AAY39801;
 XX
 XX 29-NOV-1999 (first entry)
 DT
 XX
 DE Beta-amyloid protein fragment.
 XX
 KW Beta-amyloid protein; Alzheimer's disease; amyloidosis; joint swelling;
 KW long-standing inflammation; malignancy; Familial Mediterranean Fever;
 KW multiple myeloma; plasma cell dyscrasia; long-term haemodialysis; Kuru;
 KW carpal tunnel syndrome; multiple spontaneous fracture; radiolucency;
 KW endocrine tumour; medullary carcinoma; Down's syndrome; scrapie;
 KW Creutzfeldt-Jakob disease; Gerstmann Strausler Syndrome;
 KW subacute spongiform encephalopathy; therapy.
 XX
 XX Homo sapiens.
 OS
 XX US5958883-A.
 PN
 XX 28-SEP-1999.
 PD
 XX 05-JUN-1995; 95US-0461216.
 PF
 XX 23-OCT-1992; 92US-0969734.
 PR
 XX 23-SEP-1992; 92US-0950417.
 XX
 XX (UNIW) UNIV WASHINGTON.
 PA
 XX Snow AD;
 PI
 XX WPI; 1999-561062/47.
 DR
 XX Peptides of 6-8 amino acids useful for treating or preventing
 PT amyloidosis -

XX Claim 1; Column 71; 83pp; English.
PS This sequence represents a fragment of the beta-amyloid protein. The
CC invention relates to a method for treating or preventing a form of
CC amyloidosis, including Alzheimer's disease using this sequence. The
CC compositions may be useful for treating or preventing the amyloidosis
CC associated with long-standing inflammation, various forms of malignancy
CC (including B-cell type malignancies), Familial Mediterranean Fever,
CC multiple myeloma, plasma cell dyscrasias, long-term haemodialysis, carpal
CC tunnel syndrome, joint swelling, multiple spontaneous fractures,
CC radioactivity in the wrist and hip, endocrine tumours, medullary carcinoma
CC of the thyroid, diabetes, Alzheimer's disease, Down's syndrome,
CC creutzfeldt-jakob disease, Gerstmann Strausler Syndrome, kuru, scrapie
CC and other subacute spongiform encephalopathies.
XX SQ Sequence 6 AA;

Query Match 54.5%; Score 30; DB 20; Length 6;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKL 5
Db 2 HHQKL 6
|||||

RESULT 50
AAW29090
ID AAW29090 standard; peptide; 6 AA.
XX AC AAW29090;
XX 20-JUL-1999 (first entry)
XX A-beta-binding peptide fragment conjugated to cyclosporin.
XX Cyclosporin; A-beta peptide; conjugate; neurological disease;
KW Alzheimer; multiple sclerosis; amyotrophic lateral sclerosis;
KW ALS; non-immunosuppressive; amyloid plaque formation.
XX Homo sapiens.
XX Key Location/Qualifiers
FH Modified-site 6
FT /note= "The C-terminal is condensed onto the side
FT chain of Lys(7) of the cyclosporin analog described
FT in AAW29087, AAW29088, AAW29095 and AAW29097"
XX WO9910374-A1.
XX 04-MAR-1999.
XX 25-AUG-1998; 98WO-US17544.
XX 26-AUG-1997; 97US-0057751.
XX (WISC) WISCONSIN ALUMNI RES FOUND.
XX Rich DH, Solomon ME;
PI WPI; 1999-276928/23.
XX New A-beta-binding peptide conjugates and CSA analogs - useful in
PT treatment of neurological diseases e.g. Alzheimer's disease,
PT multiple sclerosis etc.
XX Claim 5; Page 98; 129pp; English.
XX New conjugates are disclosed which are of formula A-Z, in which: A is
CC (1) a cyclosporin A analogue described in AAW29087 or (2) an FK506
CC binding peptide inhibitor; and Z is a polypeptide comprising 5 or more
CC contiguous residues of A-beta peptide. The compounds are novel chemical

CC inducers of dimerization which are non-immunosuppressive and which are
CC inhibitors of A-beta peptide aggregation and deposition in amyloid
CC plaques. The adverse consequences of amyloid plaque formation can be
CC prevented or ameliorated by sequestering the A-beta peptide in monomeric
CC form with a conjugate which links the A-beta to cyclophilin or FKBP,
CC therefore providing a mechanism to minimize the amount of free A-beta
CC available for fibril formation and deposition. The compounds can be used
CC for the treatment of Alzheimer's disease, multiple sclerosis and
CC amyotrophic lateral sclerosis.
XX SQ Sequence 6 AA;

Query Match 54.5%; Score 30; DB 20; Length 6;
Best Local Similarity 100.0%; Pred. No. 6.4e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFF 8
Db 1 QKLVFF 6
|||||

Search completed: October 29, 2002, 09:37:39
Job time : 31 secs

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:23:27 ; Search time 13 Seconds
(without alignments)
18.789 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVEFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 231628 seqs, 24425594 residues

Total number of hits satisfying chosen parameters: 231628

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : Issued_Patents_AA.*
1: /cgn2_6/ptodata/1/iaa/5A_COMB.pep.*
2: /cgn2_6/ptodata/1/iaa/5B_COMB.pep.*
3: /cgn2_6/ptodata/1/iaa/6A_COMB.pep.*
4: /cgn2_6/ptodata/1/iaa/6B_COMB.pep.*
5: /cgn2_6/ptodata/1/iaa/PCRU5_COMB.pep.*
6: /cgn2_6/ptodata/1/iaa/backfiles1.pep.*

Pred. No. is the number of results predicted by chance to have a
score greater than or equal to the score of the result being printed,
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	55	100.0	15	2	US-08-612-785B-37
2	55	100.0	17	4	US-09-264-709A-2
3	55	100.0	26	1	US-08-304-585-7
4	55	100.0	28	1	US-08-346-849-4
5	55	100.0	28	1	US-08-302-808-7
6	55	100.0	28	2	US-08-609-090-2
7	55	100.0	28	2	US-08-986-948-7
8	55	100.0	28	2	US-08-293-284A-4
9	55	100.0	28	2	US-08-461-216-2
10	55	100.0	28	4	US-09-388-890-2
11	55	100.0	28	4	US-09-388-890-3
12	55	100.0	28	4	US-09-388-890-4
13	55	100.0	28	4	US-09-388-890-5
14	55	100.0	28	4	US-09-388-890-6
15	55	100.0	28	4	US-09-388-890-7
16	55	100.0	28	4	US-09-388-890-8
17	55	100.0	28	4	US-09-388-890-13
18	55	100.0	28	4	US-09-388-890-14
19	55	100.0	28	4	US-09-284-709A-1
20	55	100.0	28	4	US-08-723-661B-2
21	55	100.0	30	2	US-08-609-090-3
22	55	100.0	32	2	US-08-609-090-4
23	55	100.0	35	1	US-08-304-585-6
24	55	100.0	35	2	US-08-612-785B-36
25	55	100.0	35	2	US-08-612-785B-38
26	55	100.0	35	2	US-08-612-785B-40
27	55	100.0	36	2	US-08-609-090-6

28	55	100.0	38	1	US-08-302-808-1	Sequence 1, Appli
29	55	100.0	38	2	US-07-737-371E-68	Sequence 68, Appl
30	55	100.0	38	2	US-08-986-948-1	Sequence 1, Appli
31	55	100.0	39	1	US-08-304-585-5	Sequence 5, Appli
32	55	100.0	39	1	US-08-302-808-2	Sequence 2, Appli
33	55	100.0	39	2	US-08-609-090-7	Sequence 7, Appli
34	55	100.0	39	2	US-08-682-245A-1	Sequence 1, Appli
35	55	100.0	39	2	US-08-986-948-2	Sequence 2, Appli
36	55	100.0	40	1	US-07-744-767A-1	Sequence 1, Appli
37	55	100.0	40	1	US-08-235-400-2	Sequence 2, Appli
38	55	100.0	40	1	US-08-476-464A-2	Sequence 2, Appli
39	55	100.0	40	1	US-08-304-585-1	Sequence 1, Appli
40	55	100.0	40	1	US-08-302-808-3	Sequence 3, Appli
41	55	100.0	40	2	US-08-433-734-1	Sequence 1, Appli
42	55	100.0	40	2	US-08-609-090-8	Sequence 8, Appli
43	55	100.0	40	2	US-07-737-371E-69	Sequence 69, Appli
44	55	100.0	40	2	US-08-682-245A-2	Sequence 2, Appli
45	55	100.0	40	2	US-08-986-948-3	Sequence 3, Appli
46	55	100.0	40	2	US-08-461-216-1	Sequence 1, Appli
47	55	100.0	40	4	US-08-959-148-1	Sequence 1, Appli
48	55	100.0	40	4	US-09-242-724-22	Sequence 22, Appli
49	55	100.0	40	4	US-08-723-661B-1	Sequence 1, Appli
50	55	100.0	40	5	PCT-US92-06700-1	Sequence 1, Appli

ALIGNMENTS

RESULT 1
US-08-612-785B-37
; Sequence 37, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Ab peptides that Modulate b-Amyloid
; TITLE OF INVENTION: Aggregation
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston.
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 37:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 15 amino acids
; TYPE: amino acid

Qy 1 HHQKLVFFAE 10
|
Db 13 HHQKLVFFAE 22

RESULT 5
US-08-302-808-7
; Sequence 7, Application US/08302808
; Patent No. 5750349
; GENERAL INFORMATION:
; APPLICANT: SUZUKI, No. 5750349uhiro
; APPLICANT: ODAKA, Asano
; APPLICANT: KITADA, Chieko
; TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
; DERIVATIVES AND USE THEREOF
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
; STREET: 130 WATER STREET
; CITY: BOSTON
; STATE: MA
; COUNTRY: USA
; ZIP: 02019
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: DOS
; SOFTWARE: FastSeq Version 1.5
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/302,808
; FILING DATE: 15-SEP-1994
; CLASSIFICATION: 435
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: PCT/JP94/00089
; FILING DATE: 24-JAN-1994
; APPLICATION NUMBER: 010132/1993
; FILING DATE: 25-JAN-1993
; APPLICATION NUMBER: 019035/1993
; FILING DATE: 05-FEB-1993
; APPLICATION NUMBER: 286985/1993
; FILING DATE: 16-NOV-1993
; APPLICATION NUMBER: 334773/1993
; FILING DATE: 28-DEC-1993
; ATTORNEY/AGENT INFORMATION:
; NAME: DAVID, RESNICK S
; REGISTRATION NUMBER: 34,235
; REFERENCE/DOCKET NUMBER: 44631
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617-523-3400
; TELEFAX: 617-523-6440
; TELEX: 200291 STRE
; INFORMATION FOR SEQ ID NO: 7:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; HYPOTHETICAL: NO
; ANTI-SENSE: NO
; FRAGMENT TYPE: N-terminal
; ORIGINAL SOURCE:
US-08-302-808-7

Query Match 100.0%; Score 55; DB 1; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
|
Db 13 HHQKLVFFAE 22

RESULT 6
US-08-609-090-2
; Sequence 2, Application US/08609090
; Patent No. 5840838
; GENERAL INFORMATION:
; APPLICANT: HENSLEY, Kenneth
; APPLICANT: BUTTERFIELD, D. A.
; APPLICANT: CARNEY, John M.
; APPLICANT: AKSENOV, Michael
; TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
; AN OLIGOPEPTIDE OR POLYPEPTIDES
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LOWE PRICE LEBLANC & BECKER
; STREET: 99 Canal Center Plaza, Suite 300
; CITY: Alexandria
; STATE: Virginia
; COUNTRY: USA
; ZIP: 22314
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/609,090
; FILING DATE: 29-FEB-1996
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Kraus, Eric J.
; REGISTRATION NUMBER: 36,190
; REFERENCE/DOCKET NUMBER: 434-059
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 703-684-1111
; TELEFAX: 703-684-1124
; INFORMATION FOR SEQ ID NO: 2:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-609-090-2

Query Match 100.0%; Score 55; DB 2; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
|
Db 13 HHQKLVFFAE 22

RESULT 7
US-08-986-948-7
; Sequence 7, Application US/08986948
; Patent No. 5955317
; GENERAL INFORMATION:
; APPLICANT: SUZUKI, No. 5955317uhiro
; APPLICANT: ODAKA, Asano
; APPLICANT: KITADA, Chieko
; TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
; DERIVATIVES AND USE THEREOF
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
; STREET: 130 WATER STREET
; CITY: BOSTON
; STATE: MA
; COUNTRY: USA
; ZIP: 02019
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette

COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSeq Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/986,948
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/302,808
FILING DATE: 15-SEP-1994
APPLICATION NUMBER: PCT/JP94/00089
FILING DATE: 24-JAN-1994
APPLICATION NUMBER: 010132/1993
FILING DATE: 25-JAN-1993
APPLICATION NUMBER: 019035/1993
FILING DATE: 05-FEB-1993
APPLICATION NUMBER: 286985/1993
FILING DATE: 16-NOV-1993
APPLICATION NUMBER: 334773/1993
FILING DATE: 28-DEC-1993
ATTORNEY/AGENT INFORMATION:
NAME: DAVID, RESNICK S
REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 7:
SEQUENCE CHARACTERISTICS:
LENGTH: 28 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
US-08-986-948-7

Query Match 100.0%; Score 55; DB 2; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVEFAE 10
| | | | | | | | | |
Db 13 HHQKLVEFAE 22

RESULT 8
US-08-293-284A-4
Sequence 4, Application US/08293284A
Patent No. 5955343
GENERAL INFORMATION:
APPLICANT: Holmes, Todd
APPLICANT: Zhang, Shuguang
APPLICANT: Rich, Alexander
APPLICANT: DiPersio, C. Michael
APPLICANT: Lockshin, Curtis
TITLE OF INVENTION: STABLE MACROSCOPIC MEMBRANES FORMED BY
TITLE OF INVENTION: SELF-ASSEMBLY OF AMPHIPHILIC PEPTIDES AND USES
TITLE OF INVENTION: THEREFOR
NUMBER OF SEQUENCES: 64
CORRESPONDENCE ADDRESS:
ADDRESSEE: HAMILTON, BROOK, SMITH & REYNOLDS, P.C.
STREET: Two Militia Drive
CITY: Lexington
STATE: Massachusetts
COUNTRY: U.S.A.
ZIP: 02173-4799
COMPUTER READABLE FORM: disk
MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/293,284A
FILING DATE: 22-AUG-1994
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 07/973,326
FILING DATE: 28-DEC-1992
ATTORNEY/AGENT INFORMATION:
NAME: Brook, David E.
REGISTRATION NUMBER: 22,592
REFERENCE/DOCKET NUMBER: MIT-6008A
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617) 861-6240
TELEFAX: (617) 861-9540
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 28 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-293-284A-4
Query Match 100.0%; Score 55; DB 2; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVEFAE 10
| | | | | | | | | |
Db 13 HHQKLVEFAE 22

RESULT 9
US-08-461-216-2
Sequence 2, Application US/08461216
Patent No. 5958883
GENERAL INFORMATION:
APPLICANT: Snow, A.D.
TITLE OF INVENTION: ANIMAL MODELS OF HUMAN AMYLOIDOSES
NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
ADDRESSEE: Christensen, O'Connor, Johnson and Kindness
STREET: 1420 Fifth Avenue, Suite 2800
CITY: Seattle
STATE: Washington
COUNTRY: USA
ZIP: 98101-2347
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette-5.25 inch, 1.2Mb storage
COMPUTER: IBM PC/386 Compatible
OPERATING SYSTEM: MS-DOS 4.01
SOFTWARE: Word for Windows-t
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/461,216
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 07/969,734
FILING DATE: October 23, 1992
APPLICATION NUMBER: 07/950,417
FILING DATE: September 23, 1992
ATTORNEY/AGENT INFORMATION:
NAME: Broderick, Thomas F.
REGISTRATION NUMBER: 31,332
REFERENCE/DOCKET NUMBER: UOFW-1-6707
TELECOMMUNICATION INFORMATION:
TELEPHONE: 1-206-682-8100; 1-206-224-0709 (direct)
TELEFAX: 1-206-224-0779
TELEX: 4938023
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:

; LENGTH: 28 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; DESCRIPTION: {SYMBOL 98 \f "Symbol"/A4(1-28)};
; DESCRIPTION: page 83, line 31
US-08-461-216-2

Query Match 100.0%; Score 55; DB 2; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 10
US-09-388-890-2
; Sequence 2, Application US/09388890
; Patent No. 6136548
; GENERAL INFORMATION:
; APPLICANT: ANDERSON, STEPHEN
; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: HOWREY & SIMON
; STREET: 1299 PENNSYLVANIA AVENUE, N.W.
; CITY: WASHINGTON
; STATE: D.C.
; COUNTRY: US
; ZIP: 20004
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/388,890
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/686,959
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: AUERBACH, JEFFREY I.
; REGISTRATION NUMBER: 32,680
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 383-7451
; TELEFAX: (202) 383-6610
; INFORMATION FOR SEQ ID NO: 2:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; HYPOTHETICAL: NO
; FRAGMENT TYPE: N-terminal
; ORIGINAL SOURCE:
; ORGANISM: HOMO SAPIENS
; IMMEDIATE SOURCE:
; CLONE: B(1-28) peptide of amyloid B protein
US-09-388-890-2

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 11
US-09-388-890-3
; Sequence 3, Application US/09388890
; Patent No. 6136548
; GENERAL INFORMATION:
; APPLICANT: ANDERSON, STEPHEN
; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: HOWREY & SIMON
; STREET: 1299 PENNSYLVANIA AVENUE, N.W.
; CITY: WASHINGTON
; STATE: D.C.
; COUNTRY: US
; ZIP: 20004
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/388,890
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/686,959
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: AUERBACH, JEFFREY I.
; REGISTRATION NUMBER: 32,680
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 383-7451
; TELEFAX: (202) 383-6610
; INFORMATION FOR SEQ ID NO: 3:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; HYPOTHETICAL: YES
; FRAGMENT TYPE: N-terminal
; ORIGINAL SOURCE:
; ORGANISM: HOMO SAPIENS
; IMMEDIATE SOURCE:
; CLONE: DIN B(1-28) peptide of amyloid B protein
US-09-388-890-3

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 12
US-09-388-890-4
; Sequence 4, Application US/09388890
; Patent No. 6136548
; GENERAL INFORMATION:
; APPLICANT: ANDERSON, STEPHEN
; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: HOWREY & SIMON
; STREET: 1299 PENNSYLVANIA AVENUE, N.W.
; CITY: WASHINGTON
; STATE: D.C.

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

COUNTRY: US
ZIP: 20004
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/388,890
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/686,959
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: AUERBACH, JEFFREY I.
REGISTRATION NUMBER: 32,680
TELEPHONE: (202) 383-7451
TELEFAX: (202) 383-6610
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
LENGTH: 28 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: YES
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
ORGANISM: HOMO SAPIENS
IMMEDIATE SOURCE:
CLONE: E3Q B(1-28) peptide of amyloid B protein
US-09-388-890-4

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 13
US-09-388-890-5
Sequence 5, Application US/09388890
Patent No. 6136548
GENERAL INFORMATION:
APPLICANT: ANDERSON, STEPHEN
TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: HOWREY & SIMON
STREET: 1299 PENNSYLVANIA AVENUE, N.W.
CITY: WASHINGTON
STATE: D.C.
COUNTRY: US
ZIP: 20004
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/388,890
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/686,959
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: AUERBACH, JEFFREY I.

REGISTRATION NUMBER: 32,680
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 383-7451
TELEFAX: (202) 383-6610
INFORMATION FOR SEQ ID NO: 5:
SEQUENCE CHARACTERISTICS:
LENGTH: 28 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: YES
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
ORGANISM: HOMO SAPIENS
IMMEDIATE SOURCE:
CLONE: RSQ B(1-28) peptide of amyloid B protein
US-09-388-890-5

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 14
US-09-388-890-6
Sequence 6, Application US/09388890
Patent No. 6136548
GENERAL INFORMATION:
APPLICANT: ANDERSON, STEPHEN
TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: HOWREY & SIMON
STREET: 1299 PENNSYLVANIA AVENUE, N.W.
CITY: WASHINGTON
STATE: D.C.
COUNTRY: US
ZIP: 20004
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/388,890
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/686,959
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: AUERBACH, JEFFREY I.
REGISTRATION NUMBER: 32,680
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 383-7451
TELEFAX: (202) 383-6610
INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
LENGTH: 28 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: YES
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
ORGANISM: HOMO SAPIENS
IMMEDIATE SOURCE:
CLONE: H6Q B(1-28) peptide of amyloid B protein

US-09-388-890-6

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
DB 13 HHQKLVFFAE 22

RESULT 15

US-09-388-890-7
; Sequence 7, Application US/09388890

; Patent No. 6136548

; GENERAL INFORMATION:

; APPLICANT: ANDERSON, STEPHEN

; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT

; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE

; NUMBER OF SEQUENCES: 14

; CORRESPONDENCE ADDRESS:

; ADDRESSEE: HOWREY & SIMON

; STREET: 1299 PENNSYLVANIA AVENUE, N.W.

; CITY: WASHINGTON

; STATE: D.C.

; COUNTRY: US

; ZIP: 20004

; COMPUTER READABLE FORM:

; MEDIUM TYPE: Floppy disk

; COMPUTER: IBM PC compatible

; OPERATING SYSTEM: PC-DOS/MS-DOS

; SOFTWARE: PatentIn Release #1.0, Version #1.25

; CURRENT APPLICATION DATA:

; APPLICATION NUMBER: US/09/388,890

; FILING DATE:

; CLASSIFICATION:

; PRIOR APPLICATION DATA:

; APPLICATION NUMBER: 08/686,959

; FILING DATE:

; ATTORNEY/AGENT INFORMATION:

; NAME: AUERBACH, JEFFREY I.

; REGISTRATION NUMBER: 32,680

; TELECOMMUNICATION INFORMATION:

; TELEPHONE: (202) 383-7451

; TELEFAX: (202) 383-6610

; INFORMATION FOR SEQ ID NO: 7:

; SEQUENCE CHARACTERISTICS:

; LENGTH: 28 amino acids

; TYPE: amino acid

; TOPOLOGY: linear

; MOLECULE TYPE: peptide

; HYPOTHETICAL: YES

; FRAGMENT TYPE: N-terminal

; ORIGINAL SOURCE:

; ORGANISM: HOMO SAPIENS

; IMMEDIATE SOURCE:

; CLONE: D7Q B(1-28) peptide of amyloid B protein

US-09-388-890-7

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
DB 13 HHQKLVFFAE 22

RESULT 16

US-09-388-890-8

; Sequence 8, Application US/09388890

; Patent No. 6136548

; GENERAL INFORMATION:

;

APPLICANT: ANDERSON, STEPHEN

TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT

OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE

NUMBER OF SEQUENCES: 14

CORRESPONDENCE ADDRESS:

ADDRESSEE: HOWREY & SIMON

STREET: 1299 PENNSYLVANIA AVENUE, N.W.

CITY: WASHINGTON

STATE: D.C.

COUNTRY: US

ZIP: 20004

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS

SOFTWARE: PatentIn Release #1.0, Version #1.25

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/09/388,890

FILING DATE:

CLASSIFICATION:

PRIOR APPLICATION DATA:

APPLICATION NUMBER: 08/686,959

FILING DATE:

ATTORNEY/AGENT INFORMATION:

NAME: AUERBACH, JEFFREY I.

REGISTRATION NUMBER: 32,680

TELECOMMUNICATION INFORMATION:

TELEPHONE: (202) 383-7451

TELEFAX: (202) 383-6610

INFORMATION FOR SEQ ID NO: 8:

SEQUENCE CHARACTERISTICS:

LENGTH: 28 amino acids

TYPE: amino acid

TOPOLOGY: linear

MOLECULE TYPE: peptide

HYPOTHETICAL: YES

FRAGMENT TYPE: N-terminal

ORIGINAL SOURCE:

ORGANISM: HOMO SAPIENS

IMMEDIATE SOURCE:

CLONE: EllQ B(1-28) peptide of amyloid B protein

US-09-388-890-8

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
DB 13 HHQKLVFFAE 22

RESULT 17

US-09-388-890-13

; Sequence 13, Application US/09388890

; Patent No. 6136548

; GENERAL INFORMATION:

; APPLICANT: ANDERSON, STEPHEN

; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT

; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE

; NUMBER OF SEQUENCES: 14

; CORRESPONDENCE ADDRESS:

; ADDRESSEE: HOWREY & SIMON

; STREET: 1299 PENNSYLVANIA AVENUE, N.W.

; CITY: WASHINGTON

; STATE: D.C.

; COUNTRY: US

; ZIP: 20004

; COMPUTER READABLE FORM:

; MEDIUM TYPE: Floppy disk

; COMPUTER: IBM PC compatible

; OPERATING SYSTEM: PC-DOS/MS-DOS

; SOFTWARE: PatentIn Release #1.0, Version #1.25

;;
;; CURRENT APPLICATION DATA:
;; APPLICATION NUMBER: US/09/388,890
;; FILING DATE:
;; CLASSIFICATION:
;; PRIOR APPLICATION DATA: 08/686,959
;; FILING DATE:
;; ATTORNEY/AGENT INFORMATION:
;; NAME: AUERBACH, JEFFREY I.
;; REGISTRATION NUMBER: 32,680
;; TELEPHONE: (202) 383-7451
;; TELEFAX: (202) 383-6610
;; INFORMATION FOR SEQ ID NO: 13:
;; SEQUENCE CHARACTERISTICS:
;; LENGTH: 28 amino acids
;; TYPE: amino acid
;; TOPOLOGY: linear
;; MOLECULE TYPE: peptide
;; HYPOTHETICAL: YES
;; FRAGMENT TYPE: N-terminal
;; ORIGINAL SOURCE:
;; ORGANISM: HOMO SAPIENS
;; IMMEDIATE SOURCE:
;; CLONE: D23Q B(1-28) peptide of amyloid B protein
US-09-388-890-13

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
| | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 18
US-09-388-890-14
; Sequence 14, Application US/09388890
; Patent No. 6136548
; GENERAL INFORMATION:
; APPLICANT: ANDERSON, STEPHEN
; TITLE OF INVENTION: METHODS FOR THE PREVENTION AND TREATMENT
; OF VASCULAR HEMORRHAGING AND ALZHEIMER'S DISEASE
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: HOWREY & SIMON
; STREET: 1299 PENNSYLVANIA AVENUE, N.W.
; CITY: WASHINGTON
; STATE: D.C.
; COUNTRY: US
; ZIP: 20004
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; OPERATING SYSTEM: IBM PC compatible
; SOFTWARE: Patent In Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/388,890
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/686,959
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: AUERBACH, JEFFREY I.
; REGISTRATION NUMBER: 32,680
; TELEPHONE: (202) 383-7451
; TELEFAX: (202) 383-6610
; INFORMATION FOR SEQ ID NO: 14:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 amino acids

;;
;; TYPE: amino acid
;; TOPOLOGY: linear
;; MOLECULE TYPE: peptide
;; HYPOTHETICAL: YES
;; FRAGMENT TYPE: N-terminal
;; ORIGINAL SOURCE:
;; ORGANISM: HOMO SAPIENS
;; IMMEDIATE SOURCE:
;; CLONE: K28Q B(1-28) peptide of amyloid B protein
US-09-388-890-14

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
| | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 19
US-09-264-709A-1
; Sequence 1, Application US/09264709A
; Patent No. 6320024
; GENERAL INFORMATION:
; APPLICANT: Roberts, Eugene
; TITLE OF INVENTION: Method for Design of Substances that Enhance Memory and
; TITLE OF INVENTION: Improve the Quality of Life
; FILE REFERENCE: 2124-310
; CURRENT APPLICATION NUMBER: US/09/264,709A
; PRIOR FILING DATE: 1999-03-09
; PRIOR APPLICATION NUMBER: 08/797,782
; PRIOR FILING DATE: 1997-02-07
; NUMBER OF SEQ ID NOS: 39
; SOFTWARE: Patent In Ver. 2.1
; SEQ ID NO 1
; LENGTH: 28
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-264-709A-1

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
| | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 20
US-08-723-661B-2
; Sequence 2, Application US/08723661B
; Patent No. 6340783
; GENERAL INFORMATION:
; APPLICANT: Alan D Snow
; TITLE OF INVENTION: Animal Models of Human Amyloidoses
; NUMBER OF SEQUENCES: 7
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Patrick M. Dwyer
; STREET: 1818 Westlake Avenue N, Suite 114
; CITY: Seattle
; STATE: WA (Washington)
; COUNTRY: United States of America
; ZIP: 98109
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette - 3.50 inch, 1.44 Mb storage
; OPERATING SYSTEM: PC-DOS (Windows 98)
; SOFTWARE: WordPerfect 5.2
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/723,661B
; FILING DATE: 31-Oct-1996

;
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/461,216
; FILING DATE: 05-Jun-1995
; APPLICATION NUMBER: 07/969,734
; FILING DATE: 23-Oct-1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: 23-Sep-1992
; ATTORNEY/AGENT INFORMATION:
; NAME: Dwyer, Patrick M.
; REGISTRATION NUMBER: 32,411
; REFERENCE/DOCKET NUMBER: PROTEO.P00C1
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (206) 343-7074
; TELEFAX: (206) 343-7085
;
; INFORMATION FOR SEQ ID NO: 2:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 28 AMINO ACIDS
; TYPE: AMINO ACID
; STRANDEDNESS: SINGLE
; TOPOLOGY: LINEAR
; MOLECULE TYPE: PEPTIDE
; DESCRIPTION: /A4 (1-28); page 83, line 31
; SEQUENCE DESCRIPTION: SEQ ID NO: 2:
US-08-723-661B-2

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00025;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22
|||||

RESULT 21
US-08-609-090-3
; Sequence 3, Application US/08609090
; Patent No. 5840838
; GENERAL INFORMATION:
; APPLICANT: HENSLEY, Kenneth
; APPLICANT: BUTTERFIELD, D. A.
; APPLICANT: CARNEY, John M.
; APPLICANT: AKSENOV, Michael
; TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
; AN OLIGOPEPTIDE OR POLYPEPTIDES
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LOWE PRICE LEBLANC & BECKER
; STREET: 99 Canal Center Plaza, Suite 300
; CITY: Alexandria
; STATE: Virginia
; COUNTRY: USA
; ZIP: 22314
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentin Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/609,090
; FILING DATE: 29-FEB-1996
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Kraus, Eric J.
; REGISTRATION NUMBER: 36,190
; REFERENCE/DOCKET NUMBER: 434-059
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 703-684-1111
; TELEFAX: 703-684-1124
; INFORMATION FOR SEQ ID NO: 3:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 30 amino acids
; TYPE: amino acid

;
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-609-090-3

Query Match 100.0%; Score 55; DB 2; Length 30;
Best Local Similarity 100.0%; Pred. No. 0.00027;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22
|||||

RESULT 22
US-08-609-090-4
; Sequence 4, Application US/08609090
; Patent No. 5840838
; GENERAL INFORMATION:
; APPLICANT: HENSLEY, Kenneth
; APPLICANT: BUTTERFIELD, D. A.
; APPLICANT: CARNEY, John M.
; APPLICANT: AKSENOV, Michael
; TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
; AN OLIGOPEPTIDE OR POLYPEPTIDES
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LOWE PRICE LEBLANC & BECKER
; STREET: 99 Canal Center Plaza, Suite 300
; CITY: Alexandria
; STATE: Virginia
; COUNTRY: USA
; ZIP: 22314
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentin Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/609,090
; FILING DATE: 29-FEB-1996
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Kraus, Eric J.
; REGISTRATION NUMBER: 36,190
; REFERENCE/DOCKET NUMBER: 434-059
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 703-684-1111
; TELEFAX: 703-684-1124
; INFORMATION FOR SEQ ID NO: 4:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 33 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-609-090-4

Query Match 100.0%; Score 55; DB 2; Length 33;
Best Local Similarity 100.0%; Pred. No. 0.0003;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22
|||||

RESULT 23
US-08-304-585-6
; Sequence 6, Application US/08304585
; Patent No. 5721106
; GENERAL INFORMATION:
; APPLICANT: Magglio, John E.

APPLICANT: Mantyh, Patrick W.
TITLE OF INVENTION: LABELLED BETA-AMYLOID PEPTIDE AND
METHODS FOR USE IN DETECTING ALZHEIMER'S DISEASE
NUMBER OF SEQUENCES: 12
CORRESPONDENCE ADDRESSES:
ADDRESSEE: Muetting, Raasch, Gebhardt & Schwappach, P.A.
STREET: P.O. Box 581415
CITY: Minneapolis
STATE: MN
COUNTRY: USA
ZIP: 55458-1415
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/304,585
FILING DATE: 12-SEP-1994
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Muetting, Ann M.
REGISTRATION NUMBER: 33,977
REFERENCE/DOCKET NUMBER: 110.00010120
TELECOMMUNICATION INFORMATION:
TELEPHONE: 612-305-1217
TELEFAX: 612-305-1228
INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
LENGTH: 35 amino acids
TYPE: amino acid
STRANDEDNESS: not relevant
TOPOLOGY: not relevant
MOLECULE TYPE: peptide
US-08-304-585-6

Query Match 100.0%; Score 55; DB 1; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00032;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 24
US-08-612-785B-36
Sequence 36, Application US/08612785B
Patent No. 5854204
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD
STREET: 28 State Street, Suite 510
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/612,785B
FILING DATE: Herewith
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995

PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 36:
SEQUENCE CHARACTERISTICS:
LENGTH: 35 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
FRAGMENT TYPE: internal
US-08-612-785B-36

Query Match 100.0%; Score 55; DB 2; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00032;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 8 HHQKLVFFAE 17

RESULT 25
US-08-612-785B-38
Sequence 38, Application US/08612785B
Patent No. 5854204
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD
STREET: 28 State Street, Suite 510
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/612,785B
FILING DATE: Herewith
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 38:

SEQUENCE CHARACTERISTICS:
; LENGTH: 35 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; FRAGMENT TYPE: internal
US-08-612-785B-38

Query Match 100.0%; Score 55; DB 2; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00032;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |
Db 13 HHQKLVFFAE 22

RESULT 26

US-08-612-785B-40
; Sequence 40, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:

APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: AB Peptides that Modulate b-Amyloid
; TITLE OF INVENTION: Aggregation
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD
STREET: 28 State Street, Suite 510
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/612.785B
FILING DATE: Herewith
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995

ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214

INFORMATION FOR SEQ ID NO: 40:
SEQUENCE CHARACTERISTICS:
LENGTH: 35 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
FRAGMENT TYPE: internal
US-08-612-785B-40

Query Match 100.0%; Score 55; DB 2; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00032;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |

Db 8 HHQKLVFFAE 17

RESULT 27

US-08-609-090-6
; Sequence 6, Application US/08609090
; Patent No. 5840838
; GENERAL INFORMATION:

APPLICANT: HENSLEY, Kenneth
APPLICANT: BUTTERFIELD, D. A.
APPLICANT: CARNEY, John M.
APPLICANT: AKSENOV, Michael
; TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
; TITLE OF INVENTION: AN OLIGOPEPTIDE OR POLYPEPTIDES
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:

ADDRESSEE: LOWE PRICE LEBLANC & BECKER
STREET: 99 Canal Center Plaza, Suite 300
CITY: Alexandria
STATE: Virginia
COUNTRY: USA
ZIP: 22314

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/609.090
FILING DATE: 29-FEB-1996
CLASSIFICATION: 530
ATTORNEY/AGENT INFORMATION:

NAME: Kraus, Eric J.
REGISTRATION NUMBER: 36,190
REFERENCE/DOCKET NUMBER: 434-059
TELECOMMUNICATION INFORMATION:
TELEPHONE: 703-684-1111
TELEFAX: 703-684-1124

INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
LENGTH: 36 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-609-090-6

Query Match 100.0%; Score 55; DB 2; Length 36;
Best Local Similarity 100.0%; Pred. No. 0.00033;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
| | | | | | | | | |

Db 13 HHQKLVFFAE 22

RESULT 28

US-08-302-808-1
; Sequence 1, Application US/08302808
; Patent No. 5750349
; GENERAL INFORMATION:

APPLICANT: SUZUKI, No. 5750349uhiro
APPLICANT: ODAKA, Asano
APPLICANT: KITADA, Chieko
; TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
; TITLE OF INVENTION: DERIVATIVES AND USE THEREOF
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:

ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
STREET: 130 WATER STREET
CITY: BOSTON
STATE: MA
COUNTRY: USA

```
;
; ZIP: 02019
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: DOS
; SOFTWARE: FastSeq Version 1.5
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/302,808
; FILING DATE: 15-SEP-1994
; CLASSIFICATION: 435
; PRIORITY APPLICATION DATA:
; APPLICATION NUMBER: PCT/JP94/00089
; FILING DATE: 24-JAN-1994
; APPLICATION NUMBER: 010132/1993
; FILING DATE: 25-JAN-1993
; APPLICATION NUMBER: 019035/1993
; FILING DATE: 05-FEB-1993
; APPLICATION NUMBER: 286985/1993
; FILING DATE: 16-NOV-1993
; APPLICATION NUMBER: 334773/1993
; FILING DATE: 28-DEC-1993
; ATTORNEY/AGENT INFORMATION:
; NAME: DAVID, RESNICK S.
; REGISTRATION NUMBER: 34,235
; REFERENCE/DOCKET NUMBER: 44631
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617-523-3400
; TELEFAX: 617-523-6440
; TELEX: 200291 STRE
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 38 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; HYPOTHETICAL: NO
; ANTI-SENSE: NO
; FRAGMENT TYPE: N-terminal
; ORIGINAL SOURCE:
; US-08-302-808-1
;
; Query Match 100.0%; Score 55; DB 1; Length 38;
; Best Local Similarity 100.0%; Pred. No. 0.00035;
; Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
;
; QY 1 HHQKLVFFAE 10
; Db 13 HHQKLVFFAE 22
;
; RESULT 29
; US-07-737-371E-68
; Sequence 68, Application US/07737371E
; Patent No. 5876948
; GENERAL INFORMATION:
; APPLICANT: Yakner, Bruce A.
; TITLE OF INVENTION: SCREENING METHODS TO IDENTIFY
; TITLE OF INVENTION: NEUROTOXIN INHIBITORS (AS AMENDED)
; NUMBER OF SEQUENCES: 77
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Fish & Richardson, P.C.
; STREET: 225 Franklin Street
; CITY: Boston
; STATE: MA
; COUNTRY: US
; ZIP: 02110-2804
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: Windows95
; SOFTWARE: FastSeq for Windows Version 2.0
; CURRENT APPLICATION DATA:
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; APPLICATION NUMBER: US/07/737,371E
; FILING DATE: 29-JUL-1991
; CLASSIFICATION: 536
; PRIORITY APPLICATION DATA:
; APPLICATION NUMBER: 07/559,172
; FILING DATE: 27-JUL-1990
; ATTORNEY/AGENT INFORMATION:
; NAME: Freeman, John W.
; REGISTRATION NUMBER: 29,066
; REFERENCE/DOCKET NUMBER: 00108/028002
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617-542-5070
; TELEFAX: 617-542-8906
; TELEX: 200154
; INFORMATION FOR SEQ ID NO: 68:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 38 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: protein
; US-07-737-371E-68
;
; Query Match 100.0%; Score 55; DB 2; Length 38;
; Best Local Similarity 100.0%; Pred. No. 0.00035;
; Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
;
; QY 1 HHQKLVFFAE 10
; Db 13 HHQKLVFFAE 22
;
; RESULT 30
; US-08-986-948-1
; Sequence 1, Application US/08986948
; Patent No. 5955317
; GENERAL INFORMATION:
; APPLICANT: SUZUKI, No. 5955317uhiro
; APPLICANT: ODAKA, Asano
; APPLICANT: KITADA, Chieko
; TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
; TITLE OF INVENTION: DERIVATIVES AND USE THEREOF
; NUMBER OF SEQUENCES: 14
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
; STREET: 130 WATER STREET
; CITY: BOSTON
; STATE: MA
; COUNTRY: USA
; ZIP: 02019
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: DOS
; SOFTWARE: FastSeq Version 1.5
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/986,948
; FILING DATE:
; CLASSIFICATION:
; PRIORITY APPLICATION DATA:
; APPLICATION NUMBER: 08/302,808
; FILING DATE: 15-SEP-1994
; APPLICATION NUMBER: PCT/JP94/00089
; FILING DATE: 24-JAN-1994
; APPLICATION NUMBER: 010132/1993
; FILING DATE: 25-JAN-1993
; APPLICATION NUMBER: 019035/1993
; FILING DATE: 05-FEB-1993
; APPLICATION NUMBER: 286985/1993
; FILING DATE: 16-NOV-1993
; APPLICATION NUMBER: 334773/1993
; FILING DATE: 28-DEC-1993
; ATTORNEY/AGENT INFORMATION:
; NAME: DAVID, RESNICK S
```


REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 38 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
US-08-986-948-1

Query Match 100.0%; Score 55; DB 2; Length 38;
Best Local Similarity 100.0%; Pred. No. 0.00035;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 31
US-08-304-585-5
Sequence 5, Application US/08304585
Patent No. 5721106
GENERAL INFORMATION:
APPLICANT: Maggio, John E.
TITLE OF INVENTION: LABELLED BETA-AMYLOID PEPTIDE AND
TITLE OF INVENTION: METHODS FOR USE IN DETECTING ALZHEIMER'S DISEASE
NUMBER OF SEQUENCES: 12
CORRESPONDENCE ADDRESS:
ADDRESSEE: Muetling, Raasch, Gebhardt & Schwappach, P.A.
STREET: P.O. Box 581415
CITY: Minneapolis
STATE: MN
COUNTRY: USA
ZIP: 55458-1415
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/304,585
FILING DATE: 12-SEP-1994
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Muetling, Ann M.
REGISTRATION NUMBER: 33,977
REFERENCE/DOCKET NUMBER: 110.00010120
TELECOMMUNICATION INFORMATION:
TELEPHONE: 612-305-1217
TELEFAX: 612-305-1228
INFORMATION FOR SEQ ID NO: 5:
SEQUENCE CHARACTERISTICS:
LENGTH: 39 amino acids
TYPE: amino acid
STRANDEDNESS: not relevant
TOPOLOGY: not relevant
MOLECULE TYPE: peptide
US-08-304-585-5

Query Match 100.0%; Score 55; DB 1; Length 39;
Best Local Similarity 100.0%; Pred. No. 0.00036;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 12 HHQKLVFFAE 21

RESULT 32
US-08-302-808-2
Sequence 2, Application US/08302808
Patent No. 5750349
GENERAL INFORMATION:
APPLICANT: SUZUKI, No. 5750349uhiro
APPLICANT: ODAKA, Asano
APPLICANT: KITADA, Chieko
TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
TITLE OF INVENTION: DERIVATIVES AND USE THEREOF
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
STREET: 130 WATER STREET
CITY: BOSTON
STATE: MA
COUNTRY: USA
ZIP: 02019
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FASTSEQ Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/302,808
FILING DATE: 15-SEP-1994
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: PCT/JP94/00089
FILING DATE: 24-JAN-1994
APPLICATION NUMBER: 010132/1993
FILING DATE: 25-JAN-1993
APPLICATION NUMBER: 019035/1993
FILING DATE: 05-FEB-1993
APPLICATION NUMBER: 286985/1993
FILING DATE: 16-NOV-1993
APPLICATION NUMBER: 334773/1993
FILING DATE: 28-DEC-1993
ATTORNEY/AGENT INFORMATION:
NAME: DAVID, RESNICK S
REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 39 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
US-08-302-808-2

Query Match 100.0%; Score 55; DB 1; Length 39;
Best Local Similarity 100.0%; Pred. No. 0.00036;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 33
US-08-609-090-7
Sequence 7, Application US/08609090
Patent No. 5840838
GENERAL INFORMATION:
APPLICANT: HENSLEY, Kenneth
APPLICANT: BUTTERFIELD, D. A.
APPLICANT: CARNEY, John M.
APPLICANT: AKSENOV, Michael
TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
TITLE OF INVENTION: AN OLIGOPEPTIDE OR POLYPEPTIDES
NUMBER OF SEQUENCES: 11
CORRESPONDENCE ADDRESS:
ADDRESSEE: LOWE PRICE LEBLANC & BECKER
STREET: 99 Canal Center Plaza, Suite 300
CITY: Alexandria
STATE: Virginia
COUNTRY: USA
ZIP: 22314
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/609,090
FILING DATE: 29-FEB-1996
CLASSIFICATION: 530
ATTORNEY/AGENT INFORMATION:
NAME: Kraus, Eric J.
REGISTRATION NUMBER: 36,190
REFERENCE/DOCKET NUMBER: 434-059
TELECOMMUNICATION INFORMATION:
TELEPHONE: 703-684-1111
TELEFAX: 703-684-1124
INFORMATION FOR SEQ ID NO: 7:
SEQUENCE CHARACTERISTICS:
LENGTH: 39 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-609-090-7

Query Match 100.0%; Score 55; DB 2; Length 39;
Best Local Similarity 100.0%; Pred. No. 0.00036;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 34
US-08-682-245A-1
Sequence 1, Application US/08682245A
Patent No. 5919631
GENERAL INFORMATION:
APPLICANT: GOVAL, SHEFALI
APPLICANT: PAUL, JOSEPH W
APPLICANT: RIEDEL, NORBERT G
APPLICANT: SAHASRABUDHE, SUDHIR
TITLE OF INVENTION: A METHOD OF DETERMINING THE DEGREE OF
TITLE OF INVENTION: AGGREGATION OF THE BAA PEPTIDE
NUMBER OF SEQUENCES: 5
CORRESPONDENCE ADDRESS:
ADDRESSEE: HOECHST MARTON ROUSSEL, INC.
STREET: 2110 E. GALBRAITH RD., P.O. BOX 156300
CITY: CINCINNATI
STATE: OHIO
COUNTRY: U.S.A.
ZIP: 45215-6300

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/682,245A
FILING DATE: 17-JUL-1996
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 60/039,414
FILING DATE: 16-AUG-1995
ATTORNEY/AGENT INFORMATION:
NAME: LENTZ, NELSEN L
REGISTRATION NUMBER: 38,537
REFERENCE/DOCKET NUMBER: HR-1257A
TELECOMMUNICATION INFORMATION:
TELEPHONE: 513-948-7369
TELEFAX: 513-948-7961 OR 4681
TELEX: 214320
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 39 amino acids
TYPE: amino acid
STRANDEDNESS:
TOPOLOGY: linear
MOLECULE TYPE: protein
US-08-682-245A-1

Query Match 100.0%; Score 55; DB 2; Length 39;
Best Local Similarity 100.0%; Pred. No. 0.00036;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 35
US-08-986-948-2
Sequence 2, Application US/08986948
Patent No. 5955317
GENERAL INFORMATION:
APPLICANT: SUZUKI, No. 5955317/uhiro
APPLICANT: ODAKA, Asano
APPLICANT: KITADA, Chieko
TITLE OF INVENTION: ANTIBODIES TO B-AMYLIDS OR THEIR
TITLE OF INVENTION: DERIVATIVES AND USE THEREOF
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
STREET: 130 WATER STREET
CITY: BOSTON
STATE: MA
COUNTRY: USA
ZIP: 02019
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSeq Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/986,948
FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/302,808
FILING DATE: 15-SEP-1994
APPLICATION NUMBER: PCT/JP94/00089
FILING DATE: 24-JAN-1994
APPLICATION NUMBER: 010132/1993
FILING DATE: 25-JAN-1993
APPLICATION NUMBER: 019035/1993

FILING DATE: 05-FEB-1993
APPLICATION NUMBER: 286985/1993
FILING DATE: 16-NOV-1993
APPLICATION NUMBER: 334773/1993
FILING DATE: 28-DEC-1993
ATTORNEY/AGENT INFORMATION:
NAME: DAVID, RESNICK S
REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 39 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
US-08-986-948-2

Query Match 100.0%; Score 55; DB 2; Length 39;
Best Local Similarity 100.0%; Pred. No. 0.00036;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 36
US-07-744-767A-1
Sequence 1, Application US/07744767A
Patent No. 5434050
GENERAL INFORMATION:
APPLICANT: Magglo, John E.
APPLICANT: Mantyh, Patrick W.
TITLE OF INVENTION: Labelled -Amyloid Peptide and Methods
TITLE OF INVENTION: for Use in Detecting Alzheimer's Disease
NUMBER OF SEQUENCES: 3
CORRESPONDENCE ADDRESS:
ADDRESSEE: Schwegman, Lundberg & Woessner, P.A.
STREET: 3500 IDS Center
CITY: Minneapolis
STATE: MN
COUNTRY: USA
ZIP: 55402
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/07744,767A
FILING DATE: 13-AUG-1991
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Muetting, Ann M.
REGISTRATION NUMBER: 33,977
REFERENCE/DOCKET NUMBER: 600.226-US-01
TELECOMMUNICATION INFORMATION:
TELEPHONE: 612-339-0331
TELEFAX: 612-339-3061
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
TOPOLOGY: linear

MOLECULE TYPE: peptide
US-07-744-767A-1
Query Match 100.0%; Score 55; DB 1; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 37
US-08-235-400-2
Sequence 2, Application US/08235400
Patent No. 5552426
GENERAL INFORMATION:
APPLICANT: Lunn, William H.
APPLICANT: Monn, James A.
APPLICANT: Zimmerman, Dennis M.
TITLE OF INVENTION: METHODS FOR TREATING A PHYSIOLOGICAL
TITLE OF INVENTION: DISORDER ASSOCIATED WITH BETA AMYLOID PEPTIDE
NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
ADDRESSEE: Eli Lilly and Company
STREET: Lilly Corporate Center/1104
CITY: Indianapolis
STATE: Indiana
COUNTRY: United States of America
ZIP: 46285
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/235,400
FILING DATE:
CLASSIFICATION: 514
ATTORNEY/AGENT INFORMATION:
NAME: Gaylo, Paul J.
REGISTRATION NUMBER: 36,808
REFERENCE/DOCKET NUMBER: X-9507
TELECOMMUNICATION INFORMATION:
TELEPHONE: (317) 276-0756
TELEFAX: (317) 276-3861
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-235-400-2

Query Match 100.0%; Score 55; DB 1; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 38
US-08-476-464A-2
Sequence 2, Application US/08476464A
Patent No. 5707821
GENERAL INFORMATION:
APPLICANT: RYDEL, RUSSELL E.
APPLICANT: DAPPEN, MICHAEL S.
TITLE OF INVENTION: THERAPEUTIC INHIBITION OF PHOSPHOLIPASE
TITLE OF INVENTION: A2 IN A-BETA PEPTIDE-MEDIATED NEURODEGENERATIVE DISEASE

NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
ADDRESSEE: TOWNSEND & TOWNSEND & CREW LLP
STREET: TWO EMBARCADERO CENTER, 8TH FLOOR
CITY: SAN FRANCISCO
STATE: CALIFORNIA
COUNTRY: U.S.A.
ZIP: 94111-3834
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/476,464A
FILING DATE: 07-JUN-1995
CLASSIFICATION: 514
ATTORNEY/AGENT INFORMATION:
NAME: STORELLA, JOHN R.
REGISTRATION NUMBER: 32,944
REFERENCE/DOCKET NUMBER: 15270-002300
TELEPHONE: (415)326-2400
TELEFAX: (415)576-0300
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS:
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-476-464A-2

Query Match 100.0%; Score 55; DB 1; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037; Indels 0;
Matches 10; Conservative 0; Mismatches 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

RESULT 39
US-08-304-585-1
Sequence 1, Application US/08304585
Patent No. 5721106
GENERAL INFORMATION:
APPLICANT: Maggio, John E.
ADDRESSEE: Martyh, Patrick W.
TITLE OF INVENTION: LABELLED BETA-AMYLOID PEPTIDE AND
METHODS FOR USE IN DETECTING ALZHEIMER'S DISEASE
NUMBER OF SEQUENCES: 12
CORRESPONDENCE ADDRESS:
ADDRESSEE: Muetting, Raasch, Gebhardt & Schwappach, P.A.
STREET: P.O. Box 581415
CITY: Minneapolis
STATE: MN
COUNTRY: USA
ZIP: 55458-1415
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/304,585
FILING DATE: 12-SEP-1994
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Muetting, Ann M.
REGISTRATION NUMBER: 33,977
REFERENCE/DOCKET NUMBER: 110,00010120
TELECOMMUNICATION INFORMATION:

TELEPHONE: 612-305-1217
TELEFAX: 612-305-1228
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS: not relevant
TOPOLOGY: not relevant
MOLECULE TYPE: peptide
US-08-304-585-1
Query Match 100.0%; Score 55; DB 1; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037; Indels 0;
Matches 10; Conservative 0; Mismatches 0; Gaps 0;
QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22
RESULT 40
US-08-302-808-3
Sequence 3, Application US/08302808
Patent No. 5750349
GENERAL INFORMATION:
APPLICANT: SUZUKI, No. 5750349uhiro
ADDRESSEE: ODAKA, Asano
APPLICANT: KITADA, Chieko
TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
DERIVATIVES AND USE THEREOF
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
STREET: 130 WATER STREET
CITY: BOSTON
STATE: MA
COUNTRY: USA
ZIP: 02019
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSeq Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/302,808
FILING DATE: 15-SEP-1994
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: PCT/JP94/00089
FILING DATE: 24-JAN-1994
APPLICATION NUMBER: 010132/1993
FILING DATE: 25-JAN-1993
APPLICATION NUMBER: 019035/1993
FILING DATE: 05-FEB-1993
APPLICATION NUMBER: 286985/1993
FILING DATE: 16-NOV-1993
APPLICATION NUMBER: 334773/1993
FILING DATE: 28-DEC-1993
ATTORNEY/AGENT INFORMATION:
NAME: DAVID, RESNICK S.
REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELECOMMUNICATION INFORMATION:
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide

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;
; HYPOTHETICAL: NO
; ANTI-SENSE: NO
; FRAGMENT TYPE: N-terminal
; ORIGINAL SOURCE:
US-08-302-808-3

Query Match      100.0%; Score 55; DB 1; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    ||||||
Db 13 HHQKLVFFAE 22

RESULT 41
US-08-433-734-1
; Sequence 1, Application US/08433734
; Patent No. 5837473
; GENERAL INFORMATION:
; APPLICANT: Maggio, John E.
; APPLICANT: Mantyh, Patrick W.
; TITLE OF INVENTION: -Amyloid Peptide and Methods
; TITLE OF INVENTION: Labelled for Use in Detecting Alzheimer's Disease
; NUMBER OF SEQUENCES: 3
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Muetting, Raasch, Gebhardt & Schwappach, P.A.
; STREET: P.O. Box 581415
; CITY: Minneapolis
; STATE: MN
; COUNTRY: USA
; ZIP: 55458-1415
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/433,734
; FILING DATE: 03-MAY-1995
; CLASSIFICATION: 435
; ATTORNEY/AGENT INFORMATION:
; NAME: Muetting, Ann M.
; REGISTRATION NUMBER: 33,977
; REFERENCE/DOCKET NUMBER: 110.00010102
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 612-305-1220
; TELEFAX: 612-305-1228
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-433-734-1

Query Match      100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    ||||||
Db 13 HHQKLVFFAE 22

RESULT 42
US-08-609-090-8
; Sequence 8, Application US/08609090
; Patent No. 5840838
; GENERAL INFORMATION:
; APPLICANT: HENSLEY, Kenneth
; APPLICANT: BUTTERFIELD, D. A.
; APPLICANT: CARNEY, John M.
```

```
;
; APPLICANT: AKSENOV, Michael
; TITLE OF INVENTION: A PROCESS FOR ENHANCING THE ACTIVITY OF
; TITLE OF INVENTION: AN OLIGOPEPTIDE OR POLYPEPTIDES
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LOWE PRICE LEBLANC & BECKER
; STREET: 99 Canal Center Plaza, Suite 300
; CITY: Alexandria
; STATE: Virginia
; COUNTRY: USA
; ZIP: 22314
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/609,090
; FILING DATE: 29-FEB-1996
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Kraus, Eric J.
; REGISTRATION NUMBER: 36,190
; REFERENCE/DOCKET NUMBER: 434-059
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 703-684-1111
; TELEFAX: 703-684-1124
; INFORMATION FOR SEQ ID NO: 8:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-609-090-8

Query Match      100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    ||||||
Db 13 HHQKLVFFAE 22

RESULT 43
US-07-737-371E-69
; Sequence 69, Application US/07737371E
; Patent No. 5876948
; GENERAL INFORMATION:
; APPLICANT: Yankner, Bruce A.
; TITLE OF INVENTION: SCREENING METHODS TO IDENTIFY
; TITLE OF INVENTION: NEUROTOXIN INHIBITORS (AS AMENDED)
; NUMBER OF SEQUENCES: 77
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Fish & Richardson, P.C.
; STREET: 225 Franklin Street
; CITY: Boston
; STATE: MA
; COUNTRY: US
; ZIP: 02110-2804
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: Windows95
; SOFTWARE: FastSeq for Windows Version 2.0
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/07/737,371E
; FILING DATE: 29-JUL-1991
; CLASSIFICATION: 536
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 07/559,172
; FILING DATE: 27-JUL-1990
```

ATTORNEY/AGENT INFORMATION:
NAME: Freeman, John W.
REGISTRATION NUMBER: 29,066
REFERENCE/DOCKET NUMBER: 00108/028002
TELEPHONE: 617-542-5070
TELEFAX: 617-542-8906
TELEX: 200154
INFORMATION FOR SEQ ID NO: 69:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: protein
US-07-737-371E-69

Query Match 100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
DB 13 HHOKLVFFAE 22

RESULT 44
US-08-682-245A-2
Sequence 2, Application US/08682245A
Patent No. 5919631
GENERAL INFORMATION:
APPLICANT: GOYAL, SHEFALI
APPLICANT: PAUL, JOSEPH W
APPLICANT: RIEDEL, NORBERT G
APPLICANT: SAHASRABUDHE, SUDHIR
TITLE OF INVENTION: A METHOD OF DETERMINING THE DEGREE OF
TITLE OF INVENTION: AGGREGATION OF THE B44 PEPTIDE
NUMBER OF SEQUENCES: 5
CORRESPONDENCE ADDRESS:
ADDRESSEE: HOECHST MARION ROUSSEL, INC.
STREET: 2110 E. GALBRAITH RD., P.O. BOX 156300
CITY: CINCINNATI
STATE: OHIO
COUNTRY: U.S.A.
ZIP: 45215-6300
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/682,245A
FILING DATE: 17-JUL-1996
CLASSIFICATION: 435

PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 60/039,414
FILING DATE: 16-AUG-1995
ATTORNEY/AGENT INFORMATION:
NAME: LENTZ, NELSEN L
REGISTRATION NUMBER: 38,537
REFERENCE/DOCKET NUMBER: HR-1257A
TELEPHONE: 513-948-7369
TELEFAX: 513-948-7961 OR 4681
TELEX: 214320
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS:
TOPOLOGY: linear
MOLECULE TYPE: protein
US-08-682-245A-2

Query Match 100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
DB 13 HHOKLVFFAE 22

RESULT 45
US-08-986-948-3
Sequence 3, Application US/08986948
Patent No. 5955317
GENERAL INFORMATION:
APPLICANT: SUZUKI, No. 5955317uhiro
APPLICANT: ODAKA, Asano
APPLICANT: KITADA, Chieko
TITLE OF INVENTION: ANTIBODIES TO B-AMYLOIDS OR THEIR
DERIVATIVES AND USE THEREOF
NUMBER OF SEQUENCES: 14
CORRESPONDENCE ADDRESS:
ADDRESSEE: DIKE, BRONSTEIN, ROBERTS & CUSHMAN
STREET: 130 WATER STREET
CITY: BOSTON
STATE: MA
COUNTRY: USA
ZIP: 02019
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSeq Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/986,948
FILING DATE:
CLASSIFICATION:

PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/302,808
FILING DATE: 15-SEP-1994
APPLICATION NUMBER: PCT/Jp94/00089
FILING DATE: 24-JAN-1994
APPLICATION NUMBER: 010132/1993
FILING DATE: 25-JAN-1993
APPLICATION NUMBER: 019035/1993
FILING DATE: 05-FEB-1993
APPLICATION NUMBER: 286985/1993
FILING DATE: 16-NOV-1993
APPLICATION NUMBER: 334773/1993
FILING DATE: 28-DEC-1993
ATTORNEY/AGENT INFORMATION:
NAME: DAVID, RESNICK S
REGISTRATION NUMBER: 34,235
REFERENCE/DOCKET NUMBER: 44631
TELEPHONE: 617-523-3400
TELEFAX: 617-523-6440
TELEX: 200291 STRE
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 40 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
ANTI-SENSE: NO
FRAGMENT TYPE: N-terminal
ORIGINAL SOURCE:
US-08-986-948-3

Query Match 100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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Qy 1 HHQKLVFFAE 10
    |||||
Db 13 HHQKLVFFAE 22

RESULT 46
US-08-461-216-1
; Sequence 1, Application US/08461216
; Patent No. 595883
; GENERAL INFORMATION:
; APPLICANT: Snow, A.D.
; TITLE OF INVENTION: ANIMAL MODELS OF HUMAN AMYLOIDOSES
; NUMBER OF SEQUENCES: 8
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Christensen, O'Connor, Johnson and Kindness
; STREET: 1420 Fifth Avenue, Suite 2800
; CITY: Seattle
; STATE: Washington
; COUNTRY: USA
; ZIP: 98101-2347
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette-5.25 inch, 1.2Mb storage
; COMPUTER: IBM PC/386 Compatible
; OPERATING SYSTEM: MS-DOS 4.01
; SOFTWARE: Word for Windows-t
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/461,216
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 07/969,734
; FILING DATE: October 23, 1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: September 23, 1992
; ATTORNEY/AGENT INFORMATION:
; NAME: Broderick, Thomas F.
; REGISTRATION NUMBER: 31,332
; REFERENCE/DOCKET NUMBER: UOFW-1-6707
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 1-206-682-8100; 1-206-224-0709 (direct)
; TELEFAX: 1-206-224-0779
; TELEX: 4938023
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; DESCRIPTION: (SYMBOL 98 \f "Symbol")/A4(1-40);
; DESCRIPTION: FIGURES 23-29
US-08-461-216-1

Query Match 100.0%; Score 55; DB 2; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    |||||
Db 13 HHQKLVFFAE 22

RESULT 47
US-08-959-148-1
; Sequence 1, Application US/08959148
; Patent No. 6172277
; GENERAL INFORMATION:
; APPLICANT: Tate, Barbara A.
; APPLICANT: Majocha, Ronald
; APPLICANT: Newton, Julie L.
; TITLE OF INVENTION: NON-TRANSGENIC ANIMAL MODEL OF ALZHEIMER'S DISEASE
; FILE REFERENCE: 04930/022001

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; CURRENT APPLICATION NUMBER: US/08/959,148
; CURRENT FILING DATE: 1997-10-28
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: FastSeq for Windows Version 3.0
; SEQ ID NO 1
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Homo sapiens
US-08-959-148-1

Query Match 100.0%; Score 55; DB 4; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    |||||
Db 13 HHQKLVFFAE 22

RESULT 48
US-09-242-724-22
; Sequence 22, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 22
; LENGTH: 40
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-242-724-22

Query Match 100.0%; Score 55; DB 4; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
    |||||
Db 13 HHQKLVFFAE 22

RESULT 49
US-08-723-661B-1
; Sequence 1, Application US/08723661B
; Patent No. 6340783
; GENERAL INFORMATION:
; APPLICANT: Alan D Snow
; TITLE OF INVENTION: Animal Models of Human Amyloidoses
; NUMBER OF SEQUENCES: 7
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Patrick M. Dwyer
; STREET: 1818 Westlake Avenue N, Suite 114
; CITY: Seattle
; STATE: WA (Washington)
; COUNTRY: United States of America
; ZIP: 98109
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette - 3.50 inch, 1.44 Mb storage
; COMPUTER: IBM PC
; OPERATING SYSTEM: PC-DOS (Windows 98)
; SOFTWARE: Wordperfect 5.2
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/723,661B
; FILING DATE: 31-Oct-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/461,216
; FILING DATE: 05-Jun-1995

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;
; APPLICATION NUMBER: 07/969,734
; FILING DATE: 23-Oct-1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: 23-Sep-1992
; ATTORNEY/AGENT INFORMATION:
; NAME: Dwyer, Patrick M.
; REGISTRATION NUMBER: 32,411
; REFERENCE/DOCKET NUMBER: PROTEO.P00C1
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (206) 343-7074
; TELEFAX: (206) 343-7085
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 AMINO ACIDS
; TYPE: AMINO ACID
; STRANDEDNESS: SINGLE
; TOPOLOGY: LINEAR
; MOLECULE TYPE: PEPTIDE
; DESCRIPTION: /A4 (1-40); FIGURES 23-29
; SEQUENCE DESCRIPTION: SEQ ID NO: 1:
US-08-723-661B-1

Query Match          100.0%; Score 55; DB 4; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

;
; NAME/KEY: amyloid peptide precursor
; LOCATION: Represents isolated internal
; LOCATION: sequence of 40 amino acid residues from
; LOCATION: the -amyloid peptide precursor
PCT-US92-06700-1

Query Match          100.0%; Score 55; DB 5; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

Search completed: October 29, 2002, 09:25:38
Job time : 14 secs

;
; APPLICATION NUMBER: 07/969,734
; FILING DATE: 23-Oct-1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: 23-Sep-1992
; ATTORNEY/AGENT INFORMATION:
; NAME: Dwyer, Patrick M.
; REGISTRATION NUMBER: 32,411
; REFERENCE/DOCKET NUMBER: PROTEO.P00C1
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (206) 343-7074
; TELEFAX: (206) 343-7085
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 AMINO ACIDS
; TYPE: AMINO ACID
; STRANDEDNESS: SINGLE
; TOPOLOGY: LINEAR
; MOLECULE TYPE: PEPTIDE
; DESCRIPTION: /A4 (1-40); FIGURES 23-29
; SEQUENCE DESCRIPTION: SEQ ID NO: 1:
US-08-723-661B-1

Query Match          100.0%; Score 55; DB 4; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

;
; NAME/KEY: amyloid peptide precursor
; LOCATION: Represents isolated internal
; LOCATION: sequence of 40 amino acid residues from
; LOCATION: the -amyloid peptide precursor
PCT-US92-06700-1

Query Match          100.0%; Score 55; DB 5; Length 40;
Best Local Similarity 100.0%; Pred. No. 0.00037;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

Search completed: October 29, 2002, 09:25:38
Job time : 14 secs

;
; APPLICATION NUMBER: 07/969,734
; FILING DATE: 23-Oct-1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: 23-Sep-1992
; ATTORNEY/AGENT INFORMATION:
; NAME: Dwyer, Patrick M.
; REGISTRATION NUMBER: 32,411
; REFERENCE/DOCKET NUMBER: PROTEO.P00C1
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (206) 343-7074
; TELEFAX: (206) 343-7085
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 40 amino acid residues
; TYPE: AMINO ACID
; TOPOLOGY: Linear
; MOLECULE TYPE: Peptide
; FRAGMENT TYPE: Internal Fragment
; ORIGINAL SOURCE: Synthetically Derived
; FEATURE:
; NAME/KEY: Internal fragment of the
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:23:27 ; Search time 16 Seconds
(without alignments)
60.056 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 283138 seqs, 96089334 residues

Total number of hits satisfying chosen parameters: 283138

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : PIR_71:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	55	100.0	42	2 PNO512	beta-amyloid prote
2	55	100.0	57	2 E60045	Alzheimer's diseas
3	55	100.0	57	2 F60045	Alzheimer's diseas
4	55	100.0	57	2 G60045	Alzheimer's diseas
5	55	100.0	57	2 D60045	Alzheimer's diseas
6	55	100.0	57	2 A60045	Alzheimer's diseas
7	55	100.0	57	2 B60045	Alzheimer's diseas
8	55	100.0	82	2 PQ0438	Alzheimer's diseas
9	55	100.0	695	1 A49795	Alzheimer's diseas
10	55	100.0	747	2 JH0773	Alzheimer's diseas
11	55	100.0	770	1 QRH0A4	Alzheimer's diseas
12	47	85.5	33	2 S23094	beta-amyloid prote
13	47	85.5	695	2 A27485	Alzheimer's diseas
14	47	85.5	695	2 S00550	Alzheimer's diseas
15	39	70.9	699	2 H64118	4-alpha-glucanotra
16	38	69.1	272	2 F70979	hypothetical prote
17	38	69.1	549	1 NU6C	glucose-6-phosphat
18	38	69.1	549	2 H91254	glucosephosphate i
19	38	69.1	549	2 D86095	glucosephosphate i
20	38	69.1	549	2 A01013	glucose-6-phosphat
21	38	69.1	550	2 B82330	glucose-6-phosphat
22	37	67.3	191	2 T04853	hypothetical prote
23	36	65.5	210	2 I58391	sarcoma amplified
24	36	65.5	535	2 S51577	transposase - rice
25	36	65.5	859	2 F69159	protoporphyrin IX
26	36	65.5	1668	1 C69224	cobalamin biosynth
27	35	63.6	297	2 T23909	hypothetical prote
28	35	63.6	446	2 T50786	nucleoid DNA-bindi
29	35	63.6	549	2 G84996	glucose-6-phosphat

30	35	63.6	552	2 T25496	hypothetical prote
31	35	63.6	751	2 D71860	probable outer mem
32	35	63.6	850	2 JC5047	ras GTPase-activat
33	35	63.6	2347	1 TVHURS	kinase-related pro
34	34	61.8	124	1 B54546	small peptidoglyca
35	34	61.8	140	2 C81176	hypothetical prote
36	34	61.8	270	2 AGI727	unknown protein hom
37	34	61.8	281	2 AGI357	unknown proteins ho
38	34	61.8	590	2 F95853	probable phospholi
39	34	61.8	635	2 H81793	hypothetical prote
40	34	61.8	763	2 S51300	probable membrane
41	34	61.8	1163	2 S07137	DNA-directed RNA p
42	34	61.8	1356	2 S51389	ROM2 protein - yea
43	34	61.8	1375	2 T18961	FAB1 protein homol
44	34	61.8	4427	2 PNO637	polyketide synthas
45	33	60.0	214	2 S39644	acetoin utilizatio
46	33	60.0	255	2 S41511	Brn-3a protein - m
47	33	60.0	258	2 D72217	conserved hypothet
48	33	60.0	325	2 A47003	cytokine receptor
49	33	60.0	334	2 T20562	hypothetical prote
50	33	60.0	336	2 S32170	phytoene synthetas

ALIGNMENTS

RESULT 1
PNO512
beta-amyloid protein - guinea pig (fragment)
C:Species: Cavia porcellus (guinea pig)
C:Date: 31-Dec-1993 #sequence_revision 31-Dec-1993 #text_change 17-Mar-1999
C:Accession: PNO512
R:Shimohigashi, Y.; Matsumoto, H.; Takano, Y.; Saito, R.; Iwata, T.; Kamiya, H.; Ohno
Biochem. Biophys. Res. Commun. 193, 624-630, 1993
A:Title: Receptor-mediated specific biological activity of a beta-amyloid protein fra
A:Reference number: PNO512; MUID:93290653
A:Accession: PNO512
A:Molecule type: protein
A:Residues: 1-42 <SH1>
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinas
C:Keywords: alternative splicing; amyloid

Query Match 100.0%; Score 55; DB 2; Length 42;
Best Local Similarity 100.0%; Pred. No. 0.00032;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22
|||||

RESULT 2
E60045
Alzheimer's disease amyloid beta/A4 protein precursor - sheep (fragment)
C:Species: Ovis sp. (sheep)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 28-Jul-1995
C:Accession: E60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in d
A:Reference number: A60045; MUID:92017079
A:Accession: E60045
A:Molecule type: mRNA
A:Residues: 1-57 <J0H>
A:Cross-references: EMBL:X56130
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinas
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10

|||||
Db 18 HHQKLVFFAE 27

RESULT 3

F60045
Alzheimer's disease amyloid beta/A4 protein precursor - pig (fragment)
C:Species: Sus scrofa domestica (domestic pig)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 13-Aug-1999
C:Accession: F60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in dog,
A:Reference number: A60045; MUID:92017079
A:Accession: F60045
A:Molecule type: mRNA
A:Residues: 1-57 <JOH>
A:Cross-references: EMBL:X56127; NID:g1895; PIDN:CAA39592.1; PID:g1896
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
Db 18 HHQKLVFFAE 27

RESULT 4

G60045
Alzheimer's disease amyloid beta/A4 protein precursor - guinea pig (fragment)
C:Species: Cavia porcellus (guinea pig)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 28-Jul-1995
C:Accession: G60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in dog,
A:Reference number: A60045; MUID:92017079
A:Accession: G60045
A:Molecule type: mRNA
A:Residues: 1-57 <JOH>
A:Cross-references: EMBL:X56126
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
Db 18 HHQKLVFFAE 27

RESULT 5

D60045
Alzheimer's disease amyloid beta/A4 protein precursor - bovine (fragment)
C:Species: Bos primigenius taurus (cattle)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 28-Jul-1995
C:Accession: D60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in dog,
A:Reference number: A60045; MUID:92017079
A:Accession: D60045
A:Molecule type: mRNA
A:Residues: 1-57 <JOH>

A:Cross-references: EMBL:X56124
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
Db 18 HHQKLVFFAE 27

RESULT 6

A60045
Alzheimer's disease amyloid beta/A4 protein precursor - dog (fragment)
C:Species: Canis lupus familiaris (dog)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 28-Jul-1995
C:Accession: A60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in d
A:Reference number: A60045; MUID:92017079
A:Accession: A60045
A:Molecule type: mRNA
A:Residues: 1-57 <JOH>
A:Cross-references: EMBL:X56125
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
Db 18 HHQKLVFFAE 27

RESULT 7

B60045
Alzheimer's disease amyloid beta/A4 protein precursor - polar bear (fragment)
C:Species: Ursus maritimus (polar bear)
C:Date: 01-Dec-1992 #sequence_revision 01-Dec-1992 #text_change 13-Aug-1999
C:Accession: B60045
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in d
A:Reference number: A60045; MUID:92017079
A:Accession: B60045
A:Molecule type: mRNA
A:Residues: 1-57 <JOH>

A:Cross-references: EMBL:X56128; NID:g2165; PIDN:CAA39593.1; PID:g2166
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; brain

Query Match 100.0%; Score 55; DB 2; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
Db 18 HHQKLVFFAE 27

RESULT 8

PQ0438
Alzheimer's disease amyloid A4 protein precursor - rabbit (fragment)
C:Species: Oryctolagus cuniculus (domestic rabbit)
C:Date: 30-Sep-1993 #sequence_revision 19-Oct-1995 #text_change 19-Oct-1995
C:Accession: PQ0438; C60045
R:Davidson, J.S.; West, R.L.; Kotikalapudi, P.; Maroun, L.E.
Biochem. Biophys. Res. Commun. 188, 905-911, 1992
A:Title: Sequence and methylation in the beta/A4 region of the rabbit amyloid precurs
A:Reference number: PQ0438; MUID:93075180
A:Accession: PQ0438
A:Molecule type: DNA

A:Residues: 1-82 <DAV>
A:Cross-references: GB:M83558; GB:M83657
R:Johnstone, E.M.; Chaney, M.O.; Norris, F.H.; Pascual, R.; Little, S.P.
Brain Res. Mol. Brain Res. 10, 299-305, 1991
A:Title: Conservation of the sequence of the Alzheimer's disease amyloid peptide in dog.
A:Reference number: A60045; MUID:92017079
A:Accession: C60045
A:Molecule type: mRNA
A:Residues: 12-68 <JOH>
A:Cross-references: EMBL:X56129
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase i
C:Keywords: alternative splicing; Alzheimer's disease; amyloid; Down's syndrome

Query Match 100.0%; Score 55; DB 2; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.00065;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
DB 29 HHQKLVFFAE 38

RESULT 9
A49795
Alzheimer's disease amyloid beta protein precursor - crab-eating macaque
C:Species: Macaca fascicularis (crab-eating macaque)
C:Date: 10-Sep-1999 #sequence_revision 10-Sep-1999 #text_change 10-Sep-1999
C:Accession: A49795
R:Podlinsky, M.B.; Tolan, D.R.; Selkoe, D.J.
Am. J. Pathol. 136, 1423-1435, 1991
A:Title: Homology of the amyloid beta protein precursor in monkey and human supports a p
A:Reference number: A49795; MUID:91273117
A:Accession: A49795
A>Status: preliminary
A:Molecule type: mRNA
A:Residues: 1-695 <POD>
A:Cross-references: GB:M58727; NID:g342062; PIDN:AAA36829.1; PID:g342063
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase i
C:Keywords: alternative splicing

Query Match 100.0%; Score 55; DB 1; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.0061;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
|||||
DB 609 HHQKLVFFAE 618

RESULT 10
JH0773
Alzheimer's disease amyloid beta protein precursor - African clawed frog
C:Species: Xenopus laevis (African clawed frog)
C:Date: 10-Jun-1993 #sequence_revision 10-Jun-1993 #text_change 13-Aug-1999
C:Accession: JH0773
R:Okado, H.; Okamoto, H.
Biochem. Biophys. Res. Commun. 189, 1561-1568, 1992
A:Title: A Xenopus homologue of the human beta-amyloid precursor protein: developmental
A:Reference number: JH0773; MUID:93129227
A:Accession: JH0773
A:Molecule type: mRNA
A:Residues: 1-747 <OKA>
A:Cross-references: GB:S52417; NID:g263150; PIDN:AAB24853.1; PID:g263151
A:Experimental source: larva
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase i
C:Keywords: alternative splicing; amyloid
F:287-337/Domain: animal Kunitz-type proteinase inhibitor homology <BPI>

Query Match 100.0%; Score 55; DB 2; Length 747;
Best Local Similarity 100.0%; Pred. No. 0.0066;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10

Db 661 HHQKLVFFAE 670
|||||
RESULT 11
QRHUA4
Alzheimer's disease amyloid beta protein precursor [validated] - human
N:Alternate names: Alzheimer's disease amyloid A4 protein; coagulation factor X1a inh
N:Contains: amyloid beta protein long, plaque form; amyloid beta protein short, vascu
protein precursor splice form APP(770)
C:Species: Homo sapiens (man)
C:Date: 30-Jun-1987 #sequence_revision 28-Jul-1995 #text_change 15-Sep-2000
C:Accession: S02260; S05194; A33260; A35486; I39452; I39451; I39453; I59562;
4668; A28593; A29302; A60805; JLO038; S06121; A50355; A59011; A38384; S29076; S38252;
R:Lemaire, H.G.; Salbaum, J.M.; Multhaup, G.; Kang, J.; Bayney, R.M.; Unterbeck, A.;
Nucleic Acids Res. 17, 517-522, 1989
A:Title: The PreA4(695) precursor protein of Alzheimer's disease A4 amyloid is encode
A:Reference number: S02260; MUID:89128427
A:Accession: S02260
A:Molecule type: DNA
A:Residues: 1-288 'V', 365-770 <LEM1>
A:Cross-references: EMBL:X13466
A:Note: alternative splice form APP(695)
R:Lemaire, H.G.
submitted to the EMBL Data Library, November 1988
A:Reference number: S05194
A:Accession: S05194
A:Molecule type: DNA
A:Residues: 1-14, 'VV', 17-288, 'V', 365-770 <LEM2>
A:Cross-references: EMBL:X13466; NID:g35598; PIDN:CAA31830.1; PID:g871360
A:Note: alternative splice form APP(695)
R:La Fauri, G.; Lahiri, D.K.; Salton, S.R.J.; Robakis, N.K.
Biochem. Biophys. Res. Commun. 159, 297-304, 1989
A:Title: Characterization of the 5'-end region and the first two exons of the beta-pr
A:Reference number: A32277; MUID:89165870
A:Accession: A32277
A:Molecule type: DNA
A:Residues: 1-75 <LAF>
A:Cross-references: GB:M24546; GB:M24547; NID:g341202; PIDN:AAC13654.1; PID:g516074
R:Johnstone, E.M.; Chaney, M.O.; Moore, R.E.; Ward, K.E.; Norris, F.H.; Little, S.P.
Biochem. Biophys. Res. Commun. 163, 1248-1255, 1989
A:Title: Alzheimer's disease amyloid peptide is encoded by two exons and shows simila
A:Reference number: A33260; MUID:89392030
A:Accession: A33260
A:Molecule type: DNA
A:Residues: 656-737 <JOH>
A:Cross-references: GB:M29270; NID:g178863; PIDN:AA51768.1; PID:g178865
R:Prelli, F.; Levy, E.; van Duinen, S.G.; Bots, G.T.A.M.; Luyendijk, W.; Frangione, B
Biochem. Biophys. Res. Commun. 170, 301-307, 1990
A:Title: Expression of a normal and variant Alzheimer's beta-protein gene in amyloid
A:Reference number: A35486; MUID:90321244
A:Accession: A35486
A:Molecule type: DNA
A:Residues: 672-710 <PRE1>
A:Note: 693-Gln was found in DNA isolated from HCHWA-D patients
R:Yoshikai, S.I.; Sasaki, H.; Doh-ura, K.; Furuya, H.; Sakaki, Y.
Gene 87, 257-263, 1990
A:Title: Genomic organization of the human amyloid beta-protein precursor gene.
A:Reference number: I39451; MUID:90236318
A:Accession: I39452
A>Status: nucleic acid sequence not shown; translation not shown; translated from GB/
A:Molecule type: DNA
A:Residues: 1-770 <YOS1>
A:Cross-references: GB:M33112; NID:g178613; PIDN:AAB59502.1; PID:g178616
A:Accession: I39451
A>Status: nucleic acid sequence not shown; translation not shown; translated from GB/
A:Molecule type: DNA
A:Residues: 1-530, 'QWLMPVIPAFWEAKVGR' <YOS2>
A:Cross-references: GB:M34875; NID:g178608; PIDN:AAB59501.1; PID:g178615
R:Yoshikai, S.I.; Sasaki, H.; Doh-ura, K.; Furuya, H.; Sakaki, Y.
Gene 102, 291-292, 1991
A:Reference number: A59020; MUID:91340168
A:Contents: annotation; erratum

A;Note: revised physical map for reference I39451
 R;Levy, E.; Carman, M.D.; Fernandez-Madrid, I.J.; Power, M.D.; Lieberburg, I.; van Duine
 Science 248, 1124-1126, 1990
 A;Title: Mutation of the Alzheimer's disease amyloid gene in hereditary cerebral hemorrh
 A;Reference number: I39453; MUID:90260663
 A;Accession: I39453
 A;Status: translated from GB/EMBL/DBDJ
 A;Molecule type: DNA
 A;Residues: 656-737 <LEV>
 A;Cross-references: GB:M37896; NID:g178618; PIDN:AAA51727.1; PID:g178620
 A;Note: a mutation with 693-Gln is presented
 R;Murrell, J.; Farlow, M.; Ghetti, B.; Benson, M.D.
 Science 254, 97-99, 1991
 A;Title: A mutation in the amyloid precursor protein associated with hereditary Alzheimer
 A;Reference number: I59562; MUID:92022553
 A;Accession: I59562
 A;Status: translated from GB/EMBL/DBDJ
 A;Molecule type: DNA
 A;Residues: 689-716, 'F', 718-737 <MUR>
 A;Cross-references: GB:S57665; NID:g236720; PIDN:AAB19991.1; PID:g236721
 R;Kamino, K.; Orr, H.T.; Payami, H.; Wijsman, E.M.; Alonso, M.E.; Pulst, S.M.; Anderson,
 arakis, S.E.; Korenberg, J.R.; Sharma, V.; Kukull, W.; Larson, E.; Heston, L.L.; Martin,
 Am. J. Hum. Genet. 51, 998-1014, 1992
 A;Title: Linkage and mutational analysis of familial Alzheimer disease kindreds for the
 A;Reference number: A44017; MUID:93035397
 A;Accession: A44017
 A;Molecule type: DNA
 A;Residues: 687-692, 'G', 694-718 <KAM1>
 A;Cross-references: GB:S45135; NID:g257377; PIDN:AAB23645.1; PID:g257378
 A;Experimental source: familial Alzheimer disease family SB
 A;Note: sequence extracted from NCBI backbone (NCBIP:115374)
 A;Accession: B44017
 A;Molecule type: DNA
 A;Residues: 687-718 <KAM2>
 A;Cross-references: GB:S45136; NID:g257379; PIDN:AAB23646.1; PID:g257380
 A;Experimental source: familial Alzheimer disease family LIT
 A;Note: Sequence extracted from NCBI backbone (NCBIP:115376)
 A;Note: This sequence has a silent mutation
 R;Kang, J.; Lemaire, H.G.; Unterbeck, A.; Salbaum, J.M.; Masters, C.L.; Grzeschik, K.H.;
 Nature 325, 733-736, 1987
 A;Title: The precursor of Alzheimer's disease amyloid A4 protein resembles a cell-surfac
 A;Reference number: A03134; MUID:87144572
 A;Accession: A03134
 A;Molecule type: mRNA
 A;Residues: 1-288, 'V', 365-770 <KAN>
 A;Cross-references: GB:Y00264; NID:g28525; PIDN:CAA68374.1; PID:g28526
 A;Note: alternative splice form APP(695)
 R;Robakis, N.K.; Ramakrishna, N.; Wolfe, G.; Wisniewski, H.M.
 Proc. Natl. Acad. Sci. U.S.A. 84, 4190-4194, 1987
 A;Title: Molecular cloning and characterization of a cDNA encoding the cerebrovascular a
 A;Reference number: A29030; MUID:87231971
 A;Accession: A29030
 A;Molecule type: mRNA
 A;Residues: 284-288, 'V', 365-646, 'E', 648-770 <ROB>
 A;Cross-references: GB:M16765; NID:g178539; PIDN:AAA51722.1; PID:g178540
 A;Note: the authors translated the codon GAG for residue 647 as Asp
 R;Goldgaber, D.; Lerman, M.I.; McBride, O.W.; Saffiotti, U.; Gajdusek, D.C.
 Science 235, 877-880, 1987
 A;Title: Characterization and chromosomal localization of a cDNA encoding brain amyloid
 A;Reference number: A47584; MUID:97120328
 A;Accession: A47584
 A;Molecule type: mRNA
 A;Residues: 674-756, 'S', 758-770 <GOL>
 A;Cross-references: GB:M15533; NID:g178706; PIDN:AAA35540.1; PID:g178707
 A;Experimental source: brain
 R;Tanzi, R.E.; Gusella, J.F.; Watkins, P.C.; Bruns, G.A.P.; St George-Hyslop, P.; Van Ke
 Science 235, 880-884, 1987
 A;Title: Amyloid beta protein gene: cDNA, mRNA distribution, and genetic linkage near th
 A;Reference number: A47585; MUID:87120329
 A;Accession: A47585
 A;Molecule type: mRNA
 A;Residues: 674-703 <TANI>
 A;Cross-references: GB:M15532; NID:g177957; PIDN:AAA51564.1; PID:g177958

R;Dyrks, T.; Weidemann, A.; Multhaup, G.; Salbaum, J.M.; Lemaire, H.G.; Kang, J.; Mue
 EMBO J. 7, 949-957, 1988
 A;Title: Identification, transmembrane orientation and biogenesis of the amyloid A4 p
 A;Reference number: S02638; MUID:88296437
 A;Accession: S02638
 A;Molecule type: mRNA
 A;Residues: 672-678 <DYR>
 R;Tanzi, R.E.; McClatchey, A.I.; Lamperti, E.D.; Villa-Komaroff, L.; Gusella, J.F.; N
 Nature 331, 528-530, 1988
 A;Title: Protease inhibitor domain encoded by an amyloid protein precursor mRNA assoc
 A;Reference number: S00707; MUID:88122640
 A;Accession: S00707
 A;Molecule type: mRNA
 A;Residues: 286-344, 'I', 365-366 <TAN2>
 A;Cross-references: EMBL:X06982; NID:g28817; PIDN:CAA30042.1; PID:g929612
 A;Experimental source: promyelocytic leukemia cell line HL60
 A;Note: alternative splice form APP(751)
 R;Ponte, P.; Gonzalez-Dewhitt, P.; Schilling, J.; Miller, J.; Hsu, D.; Greenberg, B.;
 Nature 331, 525-527, 1988
 A;Title: A new A4 amyloid mRNA contains a domain homologous to serine proteinase inh
 A;Reference number: S00925; MUID:88122639
 A;Accession: S00925
 A;Molecule type: mRNA
 A;Residues: 1-344, 'I', 365-770 <PO2>
 A;Cross-references: GB:X06989; EMBL:Y00297; NID:g28720; PIDN:CAA30050.1; PID:g28721
 A;Note: alternative splice form APP(751)
 R;Kitaguchi, N.; Takahashi, Y.; Tokushima, Y.; Shiojiri, S.; Ito, H.
 Nature 331, 530-532, 1988
 A;Title: Novel precursor of Alzheimer's disease amyloid protein shows protease inhibi
 A;Reference number: A38949; MUID:88122641
 A;Accession: A38949
 A;Molecule type: mRNA
 A;Residues: 287-367 <KIT>
 A;Cross-references: GB:X06981; NID:g28816; PIDN:CAA30041.1; PID:g929611
 A;Experimental source: glioblastoma cell line
 A;Note: alternative splice form APP(770)
 R;Vitek, M.P.; Rasool, C.G.; de Sauvage, F.; Vitek, S.M.; Bartus, R.T.; Beer, B.; Ash
 Brain Res. Mol. Brain Res. 4, 121-131, 1988
 A;Title: Absence of mutation in the beta-amyloid cDNAs cloned from the brains of thre
 A;Reference number: A30320
 A;Accession: A30320
 A;Status: not compared with conceptual translation
 A;Molecule type: mRNA
 A;Residues: 284-286, 'V', 365-770 <VIT1>
 A;Accession: B30320
 A;Status: not compared with conceptual translation
 A;Molecule type: mRNA
 A;Residues: 122-288, 'V', 365-770 <VIT2>
 A;Accession: C30320
 A;Status: not compared with conceptual translation
 A;Molecule type: mRNA
 A;Residues: 606-770 <VIT3>
 R;Zain, S.B.; Salim, M.; Chou, W.G.; Sajdel-Sulkowska, E.M.; Majocha, R.E.; Marotta,
 Proc. Natl. Acad. Sci. U.S.A. 85, 929-933, 1988
 A;Title: Molecular cloning of amyloid CDNA derived from mRNA of the Alzheimer disease
 A;Reference number: A31087; MUID:88124954
 A;Accession: A31087
 A;Molecule type: mRNA
 A;Residues: 507-770 <ZAI>
 A;Cross-references: GB:M18734; NID:g178572; PIDN:AAA51726.1; PID:g178573
 A;Note: the authors translated the codon GAA for residue 599 as Gly, ACC for residue
 8 as Val, GTG for residue 609 as Asn, AAT for residue 610 as Gly, and GGT for residue
 A;Note: the cited Genbank accession number, J03594, is not in release 101.0
 R;Masters, C.L.; Multhaup, G.; Simms, G.; Pottgiesser, J.; Martins, R.N.; Beyreuther,
 Query Watch 100.0%; Score 55; DB 1; Length 770;
 Best Local Similarity 100.0%; Pred. No. 0.0058;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 684 HHQKLVFFAE 693

```
RESULT 12
S23094
beta-amyloid protein precursor - rat
C:Species: Rattus norvegicus (Norway rat)
C:Date: 22-Nov-1993 #sequence_revision 10-Nov-1995 #text_change 03-May-1996
C:Accession: S23094
R:Kojima, S.; Omori, M.
FEBS Lett. 304, 57-60, 1992
A:Title: Two-way cleavage of beta-amyloid protein precursor by multicatalytic proteinase
A:Reference number: S23094; MUID:92316198
A:Accession: S23094
A:Molecule type: protein
A:Residues: 1-33 <KOJ>
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase i
Query Match 85.5%; Score 47; DB 2; Length 33;
Best Local Similarity 100.0%; Pred. No. 0.0099;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2 HQKLVFFAE 10
DB 19 HQKLVFFAE 27
|||||
RESULT 13
A27485
Alzheimer's disease amyloid beta/A4 protein homolog precursor - mouse
N:Alternate names: proteinase nexin II
C:Species: Mus musculus (house mouse)
C:Date: 31-Mar-1989 #sequence_revision 31-Mar-1989 #text_change 13-Aug-1999
C:Accession: A27485; S19727; I49485
R:Yamada, T.; Sakaki, H.; Furuya, H.; Miyata, T.; Goto, I.; Sakaki, Y.
Biochem. Biophys. Res. Commun. 149, 665-671, 1987
A:Title: Complementary DNA for the mouse homolog of the human amyloid beta protein precursor
A:Reference number: A27485; MUID:88106489
A:Accession: A27485
A:Molecule type: mRNA
A:Residues: 1-695 <YAM>
A:Cross-references: GB:M18373; NID:g191568; PIDN:AAA37139.1; PID:g309085
A:Experimental source: brain
R:de Strooper, B.; van Leuven, F.; van den Berghe, H.
Biochim. Biophys. Acta 1129, 141-143, 1991
A:Title: The amyloid beta protein precursor or proteinase nexin II from mouse is closer
A:Reference number: S19727; MUID:92096458
A:Accession: S19727
A:Molecule type: mRNA
A:Residues: 1-210 'G', 212-220 'S', 222-396, 'A', 398-402, 'T', 404-448, 'A', 450-695 <STR>
A:Cross-references: EMBL:X59379
R:Izumi, R.; Yamada, T.; Yoshikai, S.; Sakaki, H.; Hattori, M.; Sakaki, Y.
Gene 112, 189-195, 1992
A:Title: Positive and negative regulatory elements for the expression of the Alzheimer's
A:Reference number: I49485; MUID:92209998
A:Accession: I49485
A:Status: translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-19 <RES>
A:Cross-references: GB:D10603; NID:g220328; PIDN:BAA01456.1; PID:g220329
C:Genetics:
A:Map position: 16C3
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinase i
C:Keywords: alternative splicing; amyloid; transmembrane protein
Query Match 85.5%; Score 47; DB 2; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.24;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2 HQKLVFFAE 10
DB 610 HQKLVFFAE 618
|||||
RESULT 14
```

```
S00550
Alzheimer's disease amyloid beta protein precursor - rat
N:Alternate names: beta-A4 amyloid protein
C:Species: Rattus norvegicus (Norway rat)
C:Date: 30-Jun-1989 #sequence_revision 30-Jun-1989 #text_change 13-Aug-1999
C:Accession: S00550; A11245; A39820; S46251
R:Shivers, B.D.; Hilbich, C.; Multhaup, G.; Salbaum, M.; Beyreuther, K.; Seeburg, P.H.
EMBO J. 7, 1365-1370, 1988
A:Title: Alzheimer's disease amyloidogenic glycoprotein: expression pattern in rat br
A:Reference number: S00550; MUID:88312583
A:Accession: S00550
A:Molecule type: mRNA
A:Residues: 1-695 <SHI>
A:Cross-references: EMBL:X07648; NID:g55616; PIDN:CAA30488.1; PID:g55617
R:Schubert, D.; Schroeder, R.; LaCorbiere, M.; Saitoh, T.; Cole, G.
Science 241, 223-226, 1988
A:Title: Amyloid beta protein precursor is possibly a heparan sulfate proteoglycan co
A:Reference number: A41245; MUID:88264430
A:Accession: A41245
A:Molecule type: protein
A:Residues: 18-37 'X', 39-40, 'X', 42-44 <SCH>
A:Note: evidence for heparan sulfate attachment
R:Hesse, L.; Beher, D.; Masters, C.L.; Multhaup, G.
FEBS Lett. 349, 109-116, 1994
A:Title: The beta-A4 amyloid precursor protein binding to copper.
A:Reference number: S46251; MUID:94320627
A:Contents: annotation; copper binding sites
A:Note: rat peptides were isolated but not sequenced
R:Potempska, A.; Styles, J.; Mehta, P.; Kim, K.S.; Miller, D.L.
J. Biol. Chem. 266, 8464-8469, 1991
A:Title: Purification and tissue level of the beta-amyloid peptide precursor of rat b
A:Reference number: A39820; MUID:91217087
A:Accession: A39820
A:Status: preliminary
A:Molecule type: protein
A:Residues: 18-32 <POT>
A:Experimental source: brain
C:Comment: Deposition of amyloid protein as neurofibrillary tangles and/or plaques is
C:Superfamily: Alzheimer's disease amyloid beta protein; animal Kunitz-type proteinas
C:Keywords: alternative splicing; amyloid; glycoprotein; transmembrane protein
F:625-648/Domain: transmembrane #status predicted <TMM>
Query Match 85.5%; Score 47; DB 2; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.24;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2 HQKLVFFAE 10
DB 610 HQKLVFFAE 618
|||||
RESULT 15
H64118
4-alpha-glucanotransferase homolog - Haemophilus influenzae (strain Rd KW20)
C:Species: Haemophilus influenzae
C:Date: 18-Aug-1995 #sequence_revision 18-Aug-1995 #text_change 08-Oct-1999
C:Accession: H64118
R:Fließmann, R.D.; Adams, M.D.; White, O.; Clayton, R.A.; Kirkness, E.F.; Kerlavage
; Gocayne, J.D.; Scott, J.; Shirley, R.; Liu, L.I.; Glodek, A.; Kelley, J.M.; Weidman
; D.M.; Brandon, R.C.; Fine, L.D.; Fritchman, J.L.; Fuhrmann, J.L.; Geoghagen, N.S.M.
Science 269, 496-512, 1995
A:Title: Whole-genome random sequencing and assembly of Haemophilus influenzae Rd.
A:Reference number: A64000; MUID:95350630
A:Accession: H64118
A:Status: nucleic acid sequence not shown; translation not shown
A:Molecule type: DNA
A:Residues: 1-699 <TIGR>
A:Cross-references: GB:U32815; GB:I42023; NID:g1574818; PIDN:AAC23003.1; PID:g1574819
C:Genetics:
A:Start codon: GTG
C:Superfamily: 4-alpha-glucanotransferase
```

Query Match 70.9%; Score 39; DB 2; Length 699;
Best Local Similarity 66.7%; Pred. No. 9.5;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHOKLVFFA 9
|||: |||
Db 349 HHEKIQFFA 357

RESULT 16
F70979
hypothetical protein Rv3277 - Mycobacterium tuberculosis (strain H37RV)
C:Species: Mycobacterium tuberculosis
C>Date: 17-Jul-1998 #sequence_revision 17-Jul-1998 #text_change 22-Oct-1999
C:Accession: F70979
R: Cole, S.T.; Brosch, R.; Parkhill, J.; Garnier, T.; Churcher, C.; Harris, D.; Gordon, S.; Connor, R.T.; Davies, R.; Devlin, K.; Reltwell, T.; Gentles, S.; Hamlin, N.; Holroyd, S.; Rajandream, M.A.; Rogers, J.; Rutter, S.; Seeger, K.; Skelton, S.; Squares, S.
Nature 393, 537-544, 1998
A: Authors: Squires, R.; Sulston, J.E.; Taylor, K.; Whitehead, S.; Barrell, B.G.
A: Title: Deciphering the biology of Mycobacterium tuberculosis from the complete genome
A: Reference number: A70500; MUID: 98295987
A: Accession: F70979
A: Status: preliminary; nucleic acid sequence not shown; translation not shown
A: Molecule type: DNA
A: Residues: 1-272 <COL>
A: Cross-references: GB:292771; GB:AL123456; NID: g3242259; PIDN: CAB07080.1; PID: e306544;
A: Experimental source: strain H37RV
C: Geneticks:
A: Gene: Rv3277

Query Match 69.1%; Score 38; DB 2; Length 272;
Best Local Similarity 66.7%; Pred. No. 5.6;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHOKLVFFA 9
||: |||
Db 137 HHEALLFFA 145

RESULT 17
NUCE
Glucose-6-phosphate isomerase (EC 5.3.1.9) - Escherichia coli
N: Alternate names: phosphoglucose isomerase; phosphohexose isomerase
C: Species: Escherichia coli
C: Date: 31-Mar-1990 #sequence_revision 17-Oct-1997 #text_change 08-Sep-2000
C: Accession: H65209; J50142; S04396
R: Blattner, F.R.; Plunkett III, G.; Bloch, C.A.; Perna, N.T.; Burland, V.; Riley, M.; Cohen, A.; Rose, D.J.; Mau, B.; Shao, Y.
Science 277, 1453-1462, 1997
A: Title: The complete genome sequence of Escherichia coli K-12.
A: Reference number: A64720; MUID: 97426617
A: Accession: H65209
A: Status: preliminary; nucleic acid sequence not shown; translation not shown
A: Molecule type: DNA
A: Residues: 1-549 <BLAT>
A: Cross-references: GB:AE000476; GB:U00096; NID: g1790456; PIDN: AAC76995.1; PID: g1790457;
A: Experimental source: strain K-12, substrain MG1655
R: Froman, B.E.; Tait, R.C.; Gottlieb, L.D.
Mol. Gen. Genet. 217, 126-131, 1989
A: Title: Isolation and characterization of the phosphoglucose isomerase gene from Escherichia coli K-12.
A: Reference number: J50142; MUID: 89364675
A: Accession: J50142
A: Molecule type: DNA
A: Residues: 1-316, 'V', 318-549 <PRO>
A: Cross-references: GB: X15196; NID: g42376; PIDN: CAA33268.1; PID: g42377
A: Experimental source: strain JM101
A: Note: the authors translated the codon CAG for residue 8 as Trp
C: Comment: this enzyme catalyzes the reversible isomerization of glucose-6-phosphate and fructose-6-phosphate
C: Geneticks:
A: Gene: pgi
A: Map position: 91 min
C: Superfamily: glucose-6-phosphate isomerase

Query Match 69.1%; Score 38; DB 2; Length 549;
Best Local Similarity 66.7%; Pred. No. 12;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
||||: |||:
Db 416 HHOKLLSNEFFAQ 427

RESULT 18
H91254
glucosephosphate isomerase [imported] - Escherichia coli (strain O157:H7, substrain R
C: Species: Escherichia coli
C: Date: 18-Jul-2001 #sequence_revision 18-Jul-2001 #text_change 03-Aug-2001
C: Accession: H91254
R: Hayashi, T.; Makino, K.; Ohnishi, M.; Kurokawa, K.; Ishii, K.; Yokoyama, K.; Han, C.
gasawara, N.; Yasunaga, T.; Kuhara, S.; Shiba, T.; Hattori, M.; Shinagawa, H.
DNA Res. 8, 11-22, 2001
A: Title: Complete genome sequence of enterohemorrhagic Escherichia coli O157:H7 and
A: Reference number: A99629; MUID: 21156231; PMID: 11258796
A: Accession: H91254
A: Status: preliminary
A: Molecule type: DNA
A: Residues: 1-549 <HAY>
A: Cross-references: GB:BA000007; PIDN: BAB38431.1; PID: g13364485; GSPDB: GN00154
A: Experimental source: strain O157:H7, substrain RIMD 0509952
C: Geneticks:
A: Gene: ECs5008
C: Superfamily: glucose-6-phosphate isomerase

Query Match 69.1%; Score 38; DB 2; Length 549;
Best Local Similarity 66.7%; Pred. No. 12;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
||||: |||:
Db 416 HHOKLLSNEFFAQ 427

RESULT 19
D86095
Glucosephosphate isomerase [imported] - Escherichia coli (strain O157:H7, substrain E
C: Species: Escherichia coli
C: Date: 16-Feb-2001 #sequence_revision 16-Feb-2001 #text_change 14-Sep-2001
C: Accession: D86095
R: Perna, N.T.; Plunkett III, G.; Burland, V.; Mau, B.; Glasner, J.D.; Rose, D.J.; May
Miller, L.; Grotbeck, E.J.; Davis, N.W.; Lim, A.; Dimalanta, E.; Potamousis, K.; Apoda
Nature 409, 529-533, 2001
A: Title: Genome sequence of enterohemorrhagic Escherichia coli O157:H7.
A: Reference number: A85480; MUID: 21074935; PMID: 11206551
A: Accession: D86095
A: Status: preliminary
A: Molecule type: DNA
A: Residues: 1-549 <STO>
A: Cross-references: GB:AE005174; NID: g12518968; PIDN: AAG59224.1; GSPDB: GN00145; UWGP:
A: Experimental source: strain O157:H7, substrain EDL933
C: Geneticks:
A: Gene: pgi
C: Superfamily: glucose-6-phosphate isomerase

Query Match 69.1%; Score 38; DB 2; Length 549;
Best Local Similarity 66.7%; Pred. No. 12;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
||||: |||:
Db 416 HHOKLLSNEFFAQ 427

RESULT 20

AD1013
 glucose-6-phosphate isomerase (EC 5.3.1.9) [imported] - Salmonella enterica subsp. enterica
 C:Species: Salmonella enterica subsp. enterica serovar typhi
 A:Note: this species has also been called Salmonella typhi
 C:Date: 09-Nov-2001 #sequence_revision 09-Nov-2001 #text_change 27-Nov-2001
 C:Accession: AD1013
 R:Parkhill, J.; Dougan, G.; James, K.D.; Thomson, N.R.; Pickard, D.; Wain, J.; Churcher, T.; Connor, P.; Cronin, A.; Davis, P.; Davies, R.M.; Dowd, L.; White, N.; Farrar, S.; Moule, S.; O'Gaora, P.
 Nature 413, 848-852, 2001
 A:Authors: Parry, C.; Quail, M.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, K.; A:Title: Complete genome sequence of a multiple drug resistant Salmonella enterica serovar Paratyphi C
 A:Reference number: AB0502; PMID:11677608
 A:Accession: AD1013
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-549 <PAR>
 A:Cross-references: GB:AL513382; PIDN:CAD09205.1; PID:g16505209; GSPDB:GN00176
 C:Genetics:
 A:Gene: STY4417
 C:Superfamily: glucose-6-phosphate isomerase
 C:Keywords: intramolecular oxidoreductase; isomerase

Query Match 69.1%; Score 38; DB 2; Length 549;
 Best Local Similarity 66.7%; Pred. No. 12;
 Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
 ||||| |||
 DB 416 HHQKLLSNFFAQ 427

RESULT 21
 B82330
 glucose-6-phosphate isomerase VC0374 [imported] - Vibrio cholerae (strain N16961 serogroup O1)
 C:Species: Vibrio cholerae
 C:Date: 18-Aug-2000 #sequence_revision 20-Aug-2000 #text_change 02-Feb-2001
 C:Accession: B82330
 R:Heidelber, J.F.; Eisen, J.A.; Nelson, W.C.; Clayton, R.A.; Gwinn, M.L.; Dodson, R.J.; Churchard, D.; Ermlaeva, M.D.; Vamathevan, J.; Bass, S.; Qin, H.; Dragol, I.; Sellers, F.L.; R.R.; Mekalanos, J.J.; Venter, J.C.; Fraser, C.M.
 Nature 406, 477-483, 2000
 A:Title: DNA Sequence of both chromosomes of the cholera pathogen Vibrio cholerae.
 A:Reference number: A82035; MUID:20406833
 A:Accession: B82330
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-550 <HEI>
 A:Cross-references: GB:AE004126; GB:AE003852; NID:g9654802; PIDN:AAF93547.1; GSPDB:GN00176
 A:Experimental source: serogroup O1; strain N16961; biotype El Tor
 C:Genetics:
 A:Gene: VC0374
 A:Map position: 1
 C:Superfamily: glucose-6-phosphate isomerase

Query Match 69.1%; Score 38; DB 2; Length 550;
 Best Local Similarity 66.7%; Pred. No. 12;
 Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
 ||||| |||
 DB 417 HHQKLLSNFFAQ 428

RESULT 22
 T04853
 hypothetical protein F28A21.20 - Arabidopsis thaliana
 C:Species: Arabidopsis thaliana (mouse-ear cress)
 C:Date: 23-Apr-1999 #sequence_revision 23-Apr-1999 #text_change 04-Mar-2000
 C:Accession: T04853
 R:Bevan, M.; Mueller, M.W.; Muendlein, A.; Felber, R.; Bancroft, I.; Mewes, H.W.; Mayer, R.
 submitted to the Protein Sequence Database, February 1999
 A:Reference number: Z15387

A:Accession: T04853
 A:Molecule type: DNA
 A:Residues: 1-191 <3EV>
 A:Cross-references: EMBL:AL035526
 A:Experimental source: cultivar Columbia; BAC clone F28A21
 C:Genetics:
 A:Map position: 4
 A:Note: F28A21.20
 C:Superfamily: Arabidopsis thaliana hypothetical protein F28A21.20

Query Match 67.3%; Score 37; DB 2; Length 191;
 Best Local Similarity 60.0%; Pred. No. 6.1;
 Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 ||||| |||
 DB 86 HHQACVFFGQ 95

RESULT 23
 I58391
 sarcoma amplified sequence SAS [imported] - human
 C:Species: Homo sapiens (man)
 C:Date: 02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change 20-Jun-2000
 C:Accession: I58391
 R:Jankowski, S.A.; Mitchell, D.S.; Smith, S.H.; Trent, J.M.; Meltzer, P.S.
 Oncogene 9, 1205-1211, 1994
 A:Title: SAS, a gene amplified in human sarcomas, encodes a new member of the transmembrane protein family
 A:Reference number: I58391; MUID:94181273
 A:Accession: I58391
 A:Status: preliminary; translated from GB/EMBL/DBJ
 A:Molecule type: mRNA
 A:Residues: 1-210 <RES>
 A:Cross-references: EMBL:U01160; NID:g457936; PIDN:AAAL7782.1; PID:g457937
 C:Genetics:
 A:Gene: GDB:SAS
 A:Cross-references: GDB:I28054; OMIM:181035
 A:Map position: 12q13-12q14

Query Match 65.5%; Score 36; DB 2; Length 210;
 Best Local Similarity 75.0%; Pred. No. 11;
 Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFF 8
 ||||| |||
 DB 70 HHQVLLFF 77

RESULT 24
 S51577
 transposase - rice blast fungus
 C:Species: Magnaporthe grisea (rice blast fungus)
 C:Date: 15-Jul-1995 #sequence_revision 01-Sep-1995 #text_change 09-Sep-1997
 C:Accession: S51577
 R:Kachroo, P.; Leong, S.A.; Chattoo, B.B.
 Mol. Gen. Genet. 245, 339-348, 1994
 A:Title: Pot2, an inverted repeat transposon from the rice blast fungus Magnaporthe grisea
 A:Reference number: S51577; MUID:95115685
 A:Accession: S51577
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-535 <KAC>
 A:Cross-references: EMBL:Z33638; NID:g496853; PID:g496854

Query Match 65.5%; Score 36; DB 2; Length 535;
 Best Local Similarity 77.8%; Pred. No. 29;
 Matches 7; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQLKLVFFAE 10
 ||||| |||
 DB 80 HQLKLVFFAE 88

RESULT 25

F69159
 protoporphyrin IX magnesium chelatase (EC 4.99.1.1) - Methanobacterium thermoautotrophicum
 C:Species: Methanobacterium thermoautotrophicum
 C>Date: 05-Dec-1997 #sequence_revision 05-Dec-1997 #text_change 22-Oct-1999
 C:Accession: F69159
 R:Smith, D.R.; Doucette-Stamm, L.A.; Deloughery, C.; Lee, H.; Dubois, J.; Aldredge, T.; Qiu, D.; Spadafora, R.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Gibson, R.; Jiwani, N.; K. S.; Church, G.M.; Daniels, C.J.; Mao, J.; Rice, P.; Noelling, J.; Reeve, J.N.
 J. Bacteriol. 179, 7135-7155, 1997
 A:Title: Complete genome sequence of Methanobacterium thermoautotrophicum Delta H: functional reference genome
 A:Reference number: A69000; MUID:98037514
 A:Accession: F69159
 A:Status: preliminary; nucleic acid sequence not shown; translation not shown
 A:Molecule type: DNA
 A:Residues: 1-859 <MTH>
 A:Cross-references: GB:AE000830; GB:AE000666; NID:g2621523; PIDN:AAB84962.1; PID:g2621523
 A:Experimental source: strain Delta H
 C:Genetics:
 A:Gene: MTH456
 C:Keywords: lyase

Query Match 65.5%; Score 36; DB 2; Length 859;

Best Local Similarity 66.7%; Pred. No. 47;

Matches 6; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLVFFA 9

|||||

Db 210 HHQYLAFA 218

RESULT 26

C69224
 cobalamin biosynthesis protein N - Methanobacterium thermoautotrophicum (strain Delta H)
 C:Species: Methanobacterium thermoautotrophicum
 C>Date: 10-Sep-1999 #sequence_revision 10-Sep-1999 #text_change 21-Jul-2000
 C:Accession: C69224
 R:Smith, D.R.; Doucette-Stamm, L.A.; Deloughery, C.; Lee, H.; Dubois, J.; Aldredge, T.; Qiu, D.; Spadafora, R.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Gibson, R.; Jiwani, N.; K. S.; Church, G.M.; Daniels, C.J.; Mao, J.; Rice, P.; Noelling, J.; Reeve, J.N.
 J. Bacteriol. 179, 7135-7155, 1997
 A:Title: Complete genome sequence of Methanobacterium thermoautotrophicum Delta H: functional reference genome
 A:Reference number: A69000; MUID:98037514
 A:Accession: C69224
 A:Status: preliminary; nucleic acid sequence not shown; translation not shown
 A:Molecule type: DNA
 A:Residues: 1-1668 <MTH>
 A:Cross-references: GB:AE000868; GB:AE000666; NID:g2622025; PIDN:AAB85426.1; PID:g2622025
 A:Experimental source: strain Delta H
 C:Genetics:
 A:Gene: MTH928
 A:Start codon: GTG
 C:Superfamily: Methanobacterium thermoautotrophicum cobalamin biosynthesis protein N

Query Match 65.5%; Score 36; DB 1; Length 1668;

Best Local Similarity 66.7%; Pred. No. 94;

Matches 6; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLVFFA 9

|||||

Db 792 HHQYLAFA 800

RESULT 27

T23909
 hypothetical protein R04F11.1 - Caenorhabditis elegans
 C:Species: Caenorhabditis elegans
 C>Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 15-Oct-1999
 C:Accession: T23909
 R:Harris, B.
 submitted to the EMBL Data Library, June 1996
 A:Reference number: Z19816
 A:Accession: T23909

Query Match 63.6%; Score 35; DB 2; Length 549;

Best Local Similarity 58.3%; Pred. No. 47;

Matches 7; Conservative 2; Mismatches 1; Indels 2; Gaps 1;

A:Status: preliminary; translated from GB/EMBL/DBD

A:Molecule type: DNA

A:Residues: 1-297 <WIL>

A:Cross-references: EMBL:Z74475; PIDN:CAA98959.1; GSPDB:GN00023; CESP:R04F11.1

A:Experimental source: clone R04F11

C:Genetics:

A:Gene: CESP:R04F11.1

A:Map position: 5

A:Introns: 44/3; 82/3; 120/1; 156/1; 244/3

Query Match 63.6%; Score 35; DB 2; Length 297;

Best Local Similarity 55.6%; Pred. No. 24;

Matches 5; Conservative 3; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFFA 9

|||||

Db 157 HHKGVFFA 165

RESULT 28

T50786
 nucleoid DNA-binding protein cnd41-like protein - Arabidopsis thaliana
 N:Alternate names: protein T30N20_40
 C:Species: Arabidopsis thaliana (mouse-ear cross)
 C>Date: 21-Jul-2000 #sequence_revision 21-Jul-2000 #text_change 21-Jul-2000
 C:Accession: T50786
 R:Bevan, M.; Peters, S.A.; van Staveren, M.; Dirkse, W.; Stiekema, W.; Bancroft, I.; submitted to the Protein Sequence database, July 2000
 A:Reference number: Z25240
 A:Accession: T50786
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-446 <BEV>
 A:Cross-references: EMBL:AL365234
 A:Experimental source: cultivar Columbia; BAC clone T30N20
 C:Genetics:
 A:Map position: 5
 A:Introns: 31/3; 173/1
 A:Note: T30N20_40

Query Match 63.6%; Score 35; DB 2; Length 446;

Best Local Similarity 75.0%; Pred. No. 37;

Matches 6; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLVFF 8

|||||

Db 21 HHHHLVFF 28

RESULT 29

G84996
 glucose-6-phosphate isomerase (EC 5.3.1.9) [imported] - Buchnera sp. (strain APS)
 C:Species: Buchnera sp.
 C>Date: 02-Mar-2001 #sequence_revision 02-Mar-2001 #text_change 23-Mar-2001
 C:Accession: G84996
 R:Shigenobu, S.; Watanabe, H.; Hattori, M.; Sakaki, Y.; Ishikawa, H.
 Nature 407, 81-86, 2000
 A:Title: Genome sequence of the endocellular bacterial symbiont of aphids Buchnera sp.
 A:Reference number: A84930; MUID:20445173
 A:Accession: G84996
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-549 <STO>
 A:Cross-references: GB:AP000398; GSPDB:GN00144
 A:Experimental source: strain APS
 C:Genetics:
 A:Gene: pgi; BU573
 C:Superfamily: glucose-6-phosphate isomerase
 C:Keywords: intramolecular oxidoreductase; isomerase

Query Match

Best Local Similarity

Matches

QY 1 HHOKLV--FFAE 10
|| || - |||
Db 416 HHMKLISNFFAQ 427

RESULT 30

T25496

hypothetical protein C03G6.5 - Caenorhabditis elegans

C:Species: Caenorhabditis elegans

C:Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 15-Oct-1999

C:Accession: T25496

R:Murray, J.; Wohlmann, P.

submitted to the EMBL Data Library, April 1997

A:Description: The sequence of C. elegans cosmid C03G6.

A:Reference number: Z20042

A:Accession: T25496

A:Status: preliminary; translated from GB/EMBL/DDBJ

A:Molecule type: DNA

A:Residues: 1-552 <NR>

A:Cross-references: EMBL:U97008; PIDN:AAB52305.1; GSPDB:GN00023; CESP:C03G6.5

A:Experimental source: strain Bristol N2; clone C03G6

C:Genetics:

A:Gene: CESP:C03G6.5

A:Map position: 5

A:Introns: 28/3; 75/3; 213/3; 330/1; 393/3

Query Match

Best Local Similarity 63.6%; Score 35; DB 2; Length 552;

Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHOKLVFFFA 9

| |||: |||

Db 454 HTQKMLFFFA 462

RESULT 31

D71860

probable outer membrane protein - Helicobacter pylori (strain J99)

C:Species: Helicobacter pylori

A:Variety: strain J99

C:Date: 12-Feb-1999 #sequence_revision 12-Feb-1999 #text_change 08-Oct-1999

C:Accession: D71860

R:Alm, R.A.; Ling, L.S.L.; Moir, D.T.; King, B.L.; Brown, E.D.; Doig, P.C.; Smith, D.R.;

Ives, C.; Gibson, R.; Merberg, D.; Mills, S.D.; Jiang, Q.; Taylor, D.E.; Vovis, G.F.;

Nature 397, 176-180, 1999

A:Title: Genomic sequence comparison of two unrelated isolates of the human gastric path

A:Reference number: A71800; MUID:99120557

A:Accession: D71860

A:Status: preliminary

A:Molecule type: DNA

A:Residues: 1-751 <ARN>

A:Cross-references: GB:AE001529; GB:AE001439; NID:g4155590; PIDN:AAD06586.1; PID:g415559

A:Experimental source: strain J99

C:Genetics:

A:Gene: jhp1008

Query Match

Best Local Similarity 63.6%; Score 35; DB 2; Length 751;

Matches 7; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 2 HOKLVFFFAE 10

||| |||||

Db 26 HOKDVFFVE 34

RESULT 32

JC5047

ras GTPase-activating protein - human

C:Species: Homo sapiens (man)

C:Date: 31-Jan-1997 #sequence_revision 31-Jan-1997 #text_change 05-Nov-1999

C:Accession: JC5047

R:Kobayashi, M.; Masui, T.; Kusuda, J.; Kaneoka, Y.; Hashimoto, K.; Iwashita, S.

Gene 175, 173-177, 1996

A:Title: Human rasGTPase-activating protein (human counterpart of GAP1m): Sequence of

A:Reference number: JC5047; MUID:97074668

A:Accession: JC5047

A:Molecule type: mRNA

A:Residues: 1-850 <KOB>

A:Cross-references: DBJ:D78155; NID:g1060908; PIDN:BAAL1230.1; PID:d1011892; PID:g10

C:Comment: This protein plays a role in the regulation of cell growth and differentat

C:Genetics:

A:Gene: GAP1m

A:Map position: 3q24-26

C:Superfamily: pleckstrin repeat homology; ras-specific GAP catalytic domain homology

F:356-568/Domain: ras-specific GAP catalytic domain homology <GAP>

F:603-704/Domain: pleckstrin repeat homology <PLK>

Query Match

Best Local Similarity 63.6%; Score 35; DB 2; Length 850;

Matches 7; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHOKLVFFFA 9

|| ||| ||

Db 370 HHDKLVFFFA 378

RESULT 33

TVHURS

kinase-related protein ros-1 precursor - human

N:Alternate names: protein-tyrosine kinase mcf3 (activated ros-1)

C:Contains: protein-tyrosine kinase (EC 2.7.1.112) ros-1

C:Species: Homo sapiens (man)

C:Date: 31-Dec-1988 #sequence_revision 07-Oct-1994 #text_change 11-Jun-1999

C:Accession: A35512; A25223; A24421; A33081

R:Birchmeier, C.; O'Neill, K.; Riggs, M.; Wigler, M.

Proc. Natl. Acad. Sci. U.S.A. 87, 4799-4803, 1990

A:Title: Characterization of ROS1 cDNA from a human glioblastoma cell line.

A:Reference number: A35512; MUID:90280463

A:Accession: A35512

A:Molecule type: mRNA

A:Residues: 1-2212, 'N', 2214-2227, 'QC', 2229-2347 <BIR>

A:Cross-references: GB:M34353

A:Experimental source: glioblastoma cell line SW-1088

R:Matsumura, H.; Wang, L.H.; Shibuya, M.

Mol. Cell. Biol. 6, 3000-3004, 1986

A:Title: Human c-ros-1 gene homologous to the v-ros sequence of UR2 sarcoma virus enc

A:Reference number: A25223; MUID:87064611

A:Accession: A25223

A:Molecule type: DNA

A:Residues: 1790-2245, 'KFDSSSEFSFRCTVN' <MA2>

A:Cross-references: GB:M13368

A:Experimental source: placenta

A:Note: the differences after residue 2245 result from the authors' misinterpretation

R:Birchmeier, C.; Birnbaum, D.; Waitches, G.; Fasano, O.; Wigler, M.

Mol. Cell. Biol. 6, 3109-3116, 1986

A:Title: Characterization of an activated human ros gene.

A:Reference number: A24421; MUID:87064625

A:Accession: A24421

A:Molecule type: mRNA

A:Residues: 1854-2251, 'A', 2263-2347 <BI2>

A:Cross-references: GB:M13880; NID:g337482; PIDN:AAA36580.1; PID:g337483

A:Experimental source: tumor cells

A:Note: the mcf3 oncogene was formed by DNA rearrangement involving fusion of at least

C:Genetics:

A:Gene: GDB:ROS1

A:Cross-references: GDB:120351; OMIM:165020

A:Map position: 6q22-6q22

A:Introns: 1853/1; 1881/1; 1926/2; 1980/3; 2002/2; 2045/3; 2078/2; 2145/2; 2190/2

C:Superfamily: kinase-related protein ros; LDL receptor YWTD-containing repeat homolo

C:Keywords: ATP; autophosphorylation; glycoprotein; kinase-related transforming prote

F:1-36/Domain: signal sequence #status predicted <SIG>

F:37-2347/Product: kinase-related protein ROS1 #status predicted <MAT>

F:37-1859/Domain: extracellular #status predicted <EXT>

F:335-378/Domain: LDL receptor YWTD-containing repeat homology <YWL>

F:466-503/Domain: LDL receptor YWTD-containing repeat homology <YMA>

F:715-757/Domain: LDL receptor YWTD-containing repeat homology <YW2>
F:758-798/Domain: LDL receptor YWTD-containing repeat homology <YW3>
F:799-838/Domain: LDL receptor YWTD-containing repeat homology <YW4>
F:843-888/Domain: LDL receptor YWTD-containing repeat homology <YW5>
F:893-933/Domain: LDL receptor YWTD-containing repeat homology <YW6>
F:1532-1574/Domain: LDL receptor YWTD-containing repeat homology <YW7>
F:1860-1883/Domain: transmembrane #status predicted <TMN>
F:1884-2347/Domain: intracellular #status predicted <INT>
F:1951-1959/Region: protein kinase hmp homology <KIN>
F:1951-1959/Region: protein kinase hmp-binding motif
F:52-114,123,324,352,471,607,628,706,714,732,939,961,1015,1087,1090,1211,1272,1330,1458,
F:1960/Active site: lys #status: phosphorylated
F:2110,2114,2115/Binding site: phosphate (Tyr) (covalent) (by autophosphorylation) #stat
F:2110,2114,2115/Binding site: phosphate (Tyr) (covalent) (by autophosphorylation) #stat

Query Match 63.6% Score 35; DB 1; Length 2347;
Best Local Similarity 55.6% Pred. No. 2,1e+02;
Matches 5; Conservative 4; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQKLVFPAE 10
||:|:|:|
Db 333 HQQIVFSE 341

RESULT 34
B54546
small peptidoglycan-associated lipoprotein slp precursor - Bacillus subtilis
N;Alternate names: PAL-related lipoprotein; peptidoglycan-associated lipoprotein homolog
C;Species: Bacillus subtilis
C;Date: 03-Dec-1999 #sequence_revision 03-Dec-1999 #text_change 16-Jun-2000
C;Accession: B54546; D69708
R;Henilae, H.
FEMS Microbiol. Lett. 66, 37-41, 1991
A;Title: Sequence of a PAL-related lipoprotein from Bacillus subtilis.
A;Reference number: A54546; MUID:92038903
A;Accession: B54546
A;Molecule type: DNA
A;Residues: 1-124 <HEM>
A;Experimental source: 168 strain BBE1
A;Note: sequence extracted from NCBI backbone (NCBIN:63826, NCBIIP:63828)
R;Kunst, F.; Ogasawara, N.; Moszer, I.; Albertini, A.M.; Alloni, G.; Azevedo, V.; Berter
C.; Bron, S.; Broutillet, S.; Bruch, C.V.; Caldwell, B.; Capuano, V.; Carter, N.M.; Cho
A.; Ehrlich, S.D.; Emerson, P.T.; Entian, K.D.; Errington, J.; Faret, C.; Ferrari, E.
Nature 390, 249-256, 1997
A;Authors: Foulger, D.; Fritz, C.; Fujita, M.; Fujita, Y.; Fuma, S.; Galizzi, A.; Gallen
lech, J.; Harwood, C.R.; Henaut, A.; Hilbert, H.; Holsappel, S.; Hosono, S.; Hullo, M.F.
Koetter, P.; Konigstein, G.; Kroth, S.; Kumano, M.; Kurita, K.; Lapidus, A.; Lardinois
A;Authors: Lauber, J.; Lazarevic, V.; Lee, S.M.; Levine, A.; Liu, H.; Masuda, S.; Maue
Y, M.; Ogawa, K.; Ogiwara, A.; Oudega, B.; Park, S.H.; Parro, V.; Pohl, T.M.; Portetel
Rieger, M.; Rivolta, C.; Rocha, E.; Roche, B.; Rose, M.; Sadaie, Y.; Sato, T.; Scanlon,
A;Authors: Schleich, S.; Schroeter, R.; Scoffone, F.; Sekiguchi, J.; Sekowska, A.; Seron
akeuchi, M.; Tamakoshi, A.; Tanaka, T.; Terpstra, P.; Tognoni, A.; Tosato, V.; Uchiyama,
T.; Winters, P.; Wipat, A.; Yamamoto, H.; Yamamoto, K.; Yasumoto, K.; Yata, K.; Yoshida, K
A;Authors: Yoshikawa, H.F.; Zumbstein, E.; Yoshikawa, H.; Danchin, A.
A;Title: The complete genome sequence of the Gram-positive bacterium Bacillus subtilis.
A;Reference number: A69580; MUID:98044033
A;Accession: D69708
A;Status: nucleic acid sequence not shown; translation not shown
A;Molecule type: DNA
A;Residues: 1-124 <KUN>
A;Cross-references: GB:Z99111; GB:AL009126; NID:g2633699; PIDN:CAB13335.1; PID:g2633833
A;Experimental source: strain 168
C;Genetics:
C;Superfamily: Bacillus subtilis small peptidoglycan-associated lipoprotein slp
C;Keywords: blocked amino end; lipoprotein
F:1-18/Domain: signal sequence #status predicted <SIG>
F:19/Binding site: sn-2,3-diacylglycerol (Cys) (covalent) #status predicted
F:19/Modified site: fatty acylated amino end (Cys) (in mature form) #status predicted

Query Match 61.8% Score 34; DB 1; Length 124;
Best Local Similarity 40.0% Pred. No. 15;
Matches 4; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFPAE 10
||:|:|:|
Db 36 HHTQILFSD 45

RESULT 35
C81176
hypothetical protein NMB0648 [imported] - Neisseria meningitidis (strain MC58 serogro
C;Species: Neisseria meningitidis
C;Date: 31-Mar-2000 #sequence_revision 31-Mar-2000 #text_change 19-Jan-2001
C;Accession: C81176
R;Tettelin, H.; Saunders, N.J.; Heidelberg, J.; Jeffries, A.C.; Nelson, K.E.; Eisen,
Hickey, E.R.; Haft, D.H.; Salzberg, S.L.; White, O.; Fleischmann, R.D.; Dougherty, B.
ri, H.; Qin, H.; Vamathevan, J.; Gill, J.; Scarlato, V.; Maignani, V.; Pizza, M.
Science 287, 1809-1815, 2000
A;Authors: Grandi, G.; Sun, L.; Smith, H.O.; Fraser, C.M.; Moxon, E.R.; Rappuoli, R.;
A;Title: Complete genome sequence of Neisseria meningitidis serogroup B strain MC58.
A;Reference number: A81000; MUID:20175755
A;Accession: C81176
A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-140 <TET>
A;Cross-references: GB:AE002419; GB:AE002098; NID:g7225863; PIDN:AAF41069.1; PID:g722
A;Experimental source: serogroup B, strain MC58
C;Genetics:
C;Superfamily: Neisseria meningitidis hypothetical protein NMB0648
A;Gene: NMB0648
Query Match 61.8% Score 34; DB 2; Length 140;
Best Local Similarity 50.0% Pred. No. 18;
Matches 4; Conservative 3; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVF 8
||:|:|
Db 86 HHDVIFY 93

RESULT 36
AG1727
unknown protein homolog lin2364 [imported] - Listeria innocua (strain Clip11262)
C;Species: Listeria innocua
C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 14-Dec-2001
C;Accession: AG1727
R;Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloec
D.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi,
D.; Jones, L.M.; Karst, U.
Science 294, 849-852, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.;
Ok, C.; Schlueter, T.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehla
A;Title: Comparative genomics of Listeria species.
A;Reference number: AB1077; MUID:21537279; PMID:11679669
A;Accession: AG1727
A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-270 <GLA>
A;Cross-references: GB:AL592023; PIDN:CAC97591.1; PID:gl61414887; GSPDB:GN00178
A;Experimental source: strain Clip11262
C;Genetics:
C;Superfamily: hypothetical protein ywpJ
A;Gene: lin2364
Query Match 61.8% Score 34; DB 2; Length 270;
Best Local Similarity 60.0% Pred. No. 35;
Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 1 HHQKLVFPAE 10
||:|:|:|
Db 72 HHPRLTFPAE 81

RESULT 37
AG1357
unknown proteins homolog lmo2263 [imported] - Listeria monocytogenes (strain EGD-e)

C:Species: *Listeria monocytogenes*
C:Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
C:Accession: AG1357
R:Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloeker, R.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.; Jones, L.M.; Karst, U.
Science 294, 849-852, 2001
A:Authors: Kretz, J.; Kuhn, M.; Kunst, F.; Kurapat, G.; Madueno, E.; Maitournam, A.; Mok, C.; Schluter, T.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A.; Title: Comparative genomics of *Listeria* species.
A:Reference number: AB1077; MUID:21537279; PMID:11679669
A:Accession: AG1357
A:Status: preliminary
A:Molecule type: DNA
A:Residues: 1-281 <GLA>
A:Cross-references: GB:NC_003210; PIDN:CAD00341.1; PID:g16411733; GSPDB:GN00177
A:Experimental source: strain EGD-e
C:Genetics:
A:Gene: lmo2263

Query Match 61.8%; Score 34; DB 2; Length 281;
Best Local Similarity 60.0%; Pred. No. 37;
Matches 6; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
||| : |||
Db 83 HHPRLTTEAE 92

RESULT 38
F95853
probable phospholipase protein [imported] - *Sinorhizobium meliloti* (strain 1021) magapla
C:Species: *Sinorhizobium meliloti*
C:Date: 24-Aug-2001 #sequence_revision 24-Aug-2001 #text_change 30-Sep-2001
C:Accession: F95853
R:Finan, T.M.; Weidner, S.; Wong, K.; Buhrmester, J.; Chain, P.; Vorholter, F.J.; Hernan
Proc. Natl. Acad. Sci. U.S.A. 98, 9889-9894, 2001
A:Title: The complete sequence of the 1,683-kb pSymb megaplasmid from the N2-fixing endo
A:Reference number: A95842; MUID:21396508; PMID:11481431
A:Accession: F95853
A:Status: preliminary
A:Molecule type: DNA
A:Residues: 1-590 <KUR>
A:Cross-references: GB:AL591985; PIDN:CAC48494.1; PID:g15139966; GSPDB:GN00167
A:Experimental source: strain 1021, megaplasmid pSymb
R:Galibert, F.; Finan, T.M.; Long, S.R.; Puhler, A.; Abola, P.; Ampe, F.; Barloy-Hubler, J.
pela, D.; Chain, P.; Cowie, A.; Davis, R.W.; Dreano, S.; Federspiel, N.A.; Fisher, R.F.;
L.; Hyman, R.W.; Jones, T.
Science 293, 668-672, 2001
A:Authors: Kahn, D.; Kahn, M.L.; Kalman, S.; Keating, D.H.; Kiss, E.; Komp, C.; Lelaure,
hebaullt, P.; Vandenbol, M.; Vorholter, F.J.; Weidner, S.; Wells, D.H.; Wong, K.; Yeh, K.
A:Title: The composite genome of the legume symbiont *Sinorhizobium meliloti*.
A:Reference number: A96039; MUID:21368234; PMID:11474104
A:Contents: annotation
C:Genetics:
A:Gene: smb20094
A:Genome: plasmid

Query Match 61.8%; Score 34; DB 2; Length 590;
Best Local Similarity 100.0%; Pred. No. 79;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLV 6
|||||
Db 208 HHQKLV 213

RESULT 39
H81793
hypothetical protein NMA2205 [imported] - *Neisseria meningitidis* (strain Z2491 serogroup
C:Species: *Neisseria meningitidis*
C:Date: 05-May-2000 #sequence_revision 05-May-2000 #text_change 02-Feb-2001
C:Accession: H81793

R:Parkhill, J.; Achtman, M.; James, K.D.; Bentley, S.D.; Churcher, C.; Klee, S.R.; Mo
; Holroyd, S.; Jagels, K.; Leather, S.; Moule, S.; Mungall, K.; Quail, M.A.; Rajandre
Nature 404, 502-506, 2000
A:Title: Complete DNA sequence of a serogroup A strain of *Neisseria meningitidis* Z2491
A:Reference number: AB1775; MUID:20222556
A:Accession: H81793
A:Status: preliminary
A:Molecule type: DNA
A:Residues: 1-635 <PAR>
A:Cross-references: GB:AL162758; GB:AL157959; NID:g7380672; PIDN:CAB85416.1; PID:g738
A:Experimental source: serogroup A, strain Z2491
C:Genetics:
A:Gene: NMA2205

Query Match 61.8%; Score 34; DB 2; Length 635;
Best Local Similarity 62.5%; Pred. No. 86;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

OY 1 HHQKLVFF 8
||| : |||
Db 387 HHQDLIWF 394

RESULT 40
SS1300
Probable membrane protein YNL311c - yeast (*Saccharomyces cerevisiae*)
N:Alternate names: hypothetical protein N0376
C:Species: *Saccharomyces cerevisiae*
C:Date: 23-Feb-1995 #sequence_revision 12-May-1995 #text_change 23-Mar-2001
C:Accession: SS1300; S59569; S63292; S63284
R:Nicaud, J.J.
submitted to the EMBL Data Library, January 1995
A:Description: Sequence analysis of a 13.9 Kb fragment of yeast chromosome XIV identifi
A:Reference number: SS1285
A:Accession: SS1300
A:Molecule type: DNA
A:Residues: 1-763 <NIC>
A:Cross-references: EMBL:Z46259; NID:g633655; PID:g633671
R:Matfahl, M.; Nicaud, J.M.; Levesque, H.; Gaillardin, C.
Yeast 11, 1077-1085, 1995
A:Title: Sequencing analysis of a 24.7 kb fragment of yeast chromosome XIV identifies
A:Reference number: S59562; MUID:96076632
A:Accession: S59569
A:Status: nucleic acid sequence not shown; translation not shown
A:Molecule type: DNA
A:Residues: 1-763 <NAF>
A:Cross-references: EMBL:Z46259; NID:g633655; PIDN:CAA86384.1; PID:g633671
A:Note: the nucleotide sequence was submitted to the EMBL Data Library, January 1995
R:Matfahl, M.; Nicaud, J.M.; Levesque, H.; Gaillardin, C.
submitted to the Protein Sequence Database, April 1996
A:Reference number: S63287
A:Accession: S63292
A:Molecule type: DNA
A:Residues: 1-763 <NAW>
A:Cross-references: EMBL:Z71587; NID:g1302414; PID:g1302415; MIPS:YNL311c
A:Experimental source: strain S288C
R:Maurer, C.T.C.; Urbanus, J.H.M.; Planta, R.J.
submitted to the Protein Sequence Database, April 1996
A:Reference number: S63266
A:Accession: S63284
A:Molecule type: DNA
A:Residues: 148-763 <NAU>
A:Cross-references: EMBL:Z71587; MIPS:YNL311c
A:Experimental source: strain S288C
C:Genetics:
A:Map position: 14L
C:Superfamily: *Saccharomyces cerevisiae* probable membrane protein YNL311c
C:Keywords: transmembrane protein
F:64-80/Domain: transmembrane #status predicted <TM>

Query Match 61.8%; Score 34; DB 2; Length 763;
Best Local Similarity 100.0%; Pred. No. 1e+02;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
|||||
Db 323 HHQKLV 328

RESULT 41
S07137
DNA-directed RNA polymerase (EC 2.7.7.6) beta'-2 chain - garden pea chloroplast (fragment)
C:Species: Chloroplast Pisum sativum (garden pea)
C:Date: 31-Dec-1990 #sequence_revision 31-Dec-1990 #text_change 18-Jun-1999
C:Accession: S07137
R:Cozens, A.L.; Walker, J.E.
Biochem. J. 236, 453-460, 1986
A:Title: Pea chloroplast DNA encodes homologues of Escherichia coli ribosomal subunit S2
A:Reference number: S07137; MUID:86323089
A:Accession: S07137
A:Molecule type: DNA
A:Residues: 1-1163 <COZ>
A:Cross-references: EMBL:X03912; NID:g12137; PIDN:CAA27545.1; PID:g8293325
C:Genetics:
A:Gene: rpoC2
A:Genome: chloroplast
A:Superfamily: chloroplast DNA-directed RNA polymerase beta'-2 chain
C:Keywords: chloroplast; nucleotidyltransferase; transcription

Query Match 61.8%; Score 34; DB 2; Length 1163;
Best Local Similarity 66.7%; Pred. No. 1.6e+02;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFFA 9
||:|:|
Db 1149 HHRKLLDFA 1157

RESULT 42
S51389
ROM2 protein - yeast (Saccharomyces cerevisiae)
N:Alternate names: protein L8039.3; protein YLR371w
C:Species: Saccharomyces cerevisiae
C:Date: 23-Feb-1995 #sequence_revision 12-May-1995 #text_change 05-Nov-1999
C:Accession: S51389
R:Du, Z.
submitted to the EMBL Data Library, December 1994
A:Description: The sequence of S. cerevisiae cosmid 8039.
A:Reference number: S51377
A:Accession: S51389
A:Molecule type: DNA
A:Residues: 1-1356 <DUZ>
A:Cross-references: EMBL:U19103; NID:g609404; PID:g609407; GSPDB:GN00012; MIPS:YLR371w
C:Genetics:
A:Gene: ROM2; MIPS:YLR371w
A:Map position: 12R
C:Superfamily: CDC24 homology
F:659-846/Domain: CDC24 homology <CD24>

Query Match 61.8%; Score 34; DB 2; Length 1356;
Best Local Similarity 50.0%; Pred. No. 1.9e+02;
Matches 7; Conservative 3; Mismatches 0; Indels 4; Gaps 1;

QY 1 HHQKLV----FFAE 10
||:|:|
Db 1131 HHRKLVHVSFFAE 1144

RESULT 43
T18961
FAB1 protein homolog VF11C1L.1 - Caenorhabditis elegans
C:Species: Caenorhabditis elegans
C:Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 18-Feb-2000
C:Accession: T18961; T26005
R:Lloyd, C.
submitted to the EMBL Data Library, November 1995

A:Reference number: Z19052
A:Accession: T18961
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-1375 <WIL>
A:Cross-references: EMBL:Z67879; PIDN:CAA91791.1; GSPDB:GN00028; CESP:VF11C1L.1
A:Experimental source: clone C05E7
R:Mortimore, B.
submitted to the EMBL Data Library, June 1998
A:Reference number: Z20126
A:Accession: T26005
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-1375 <W12>
A:Cross-references: EMBL:AL023817; PIDN:CAAL9436.1; GSPDB:GN00028; CESP:VF11C1L.1
A:Experimental source: clone VF11C1L
C:Genetics:
A:Gene: CESP:VF11C1L.1
A:Map position: X
A:Introns: 109/1; 172/2; 198/3; 396/3; 561/3; 592/2; 647/1; 789/2; 859/3; 1104/3; 123

Query Match 61.8%; Score 34; DB 2; Length 1375;
Best Local Similarity 75.0%; Pred. No. 1.9e+02;
Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HOKLVFFA 9
|:|:|
Db 840 HEKLYFFA 847

RESULT 44
PN0637
polyketide synthase pksL - Bacillus subtilis
C:Species: Bacillus subtilis
C:Date: 19-May-1994 #sequence_revision 06-Feb-1995 #text_change 03-Nov-2000
C:Accession: S25021; PN0637; B69679
R:Scotti, C.; Piatti, M.; Curzoni, A.; Tognoni, A.; Grandi, G.; Galizzi, A.; Albertin
submitted to the EMBL Data Library, July 1992
A:Description: A Bacillus subtilis large ORF coding for a polypeptide highly similar
A:Reference number: S25021
A:Accession: S25021
A:Status: preliminary
A:Molecule type: DNA
A:Residues: 1-4427 <SCO>
A:Cross-references: EMBL:Z14098; NID:g40057; PIDN:CAA78479.1; PID:g40058
R:Scotti, C.; Piatti, M.; Curzoni, A.; Perani, P.; Tognoni, A.; Grandi, G.; Galizzi, A.
Gene 130, 65-71, 1993
A:Title: A Bacillus subtilis large ORF coding for a polypeptide highly similar to pol
A:Reference number: PN0637; MUID:93345824
A:Accession: PN0637
A:Status: nucleic acid sequence not shown
A:Molecule type: DNA
A:Residues: 184-282; 382-850; 926-1115; 1409-1648; 1665-1761; 1876-2344; 2469-2560; 2609-270
R:Kunst, F.; Ogasawara, N.; Moszer, I.; Albertini, A.M.; Alloni, G.; Azevedo, V.; Ber
C.; Bron, S.; Brouillet, S.; Brusch, C.V.; Caldwell, B.; Capuano, V.; Carter, N.M.;
A.; Ehrlich, S.D.; Emmerson, P.T.; Entlan, K.D.; Errington, J.; Fabret, C.; Ferrari,
Nature 390, 249-256, 1997
A:Authors: Foulger, D.; Fritz, C.; Fujita, M.; Fujita, Y.; Fuma, S.; Galizzi, A.; Gal
iech, J.; Harwood, C.R.; Henaut, A.; Hilbert, H.; Holsappel, S.; Hosono, S.; Hullo, M
Koetter, P.; Konigstein, G.; Krogh, S.; Kumano, M.; Kurita, K.; Lapidus, A.; Lardino
A:Authors: Lauber, J.; Lazarevic, V.; Lee, S.M.; Levine, A.; Liu, H.; Masuda, S.; Mau
Y, M.; Odawa, K.; Ogiwara, A.; Oudega, B.; Park, S.H.; Parro, V.; Pohl, T.M.; Portete
Rieger, M.; Rivolta, C.; Rocha, E.; Roche, B.; Rose, M.; Sadate, Y.; Sato, T.; Scani
A:Authors: Schleich, S.; Schroeter, R.; Scoffone, F.; Sekiguchi, J.; Sekowska, A.; Se
akeuchi, M.; Tamakoshi, A.; Tanaka, T.; Terpstra, P.; Tognoni, A.; Tosato, V.; Uchiya
T.; Winters, P.; Wipat, A.; Yamamoto, H.; Yamane, K.; Yasumoto, K.; Yata, K.; Yoshida
A:Authors: Yoshikawa, H.F.; Zumstein, E.; Yoshikawa, H.; Danchin, A.
A:Title: The complete genome sequence of the Gram-positive bacterium Bacillus subtili
A:Reference number: A69580; MUID:98044033
A:Accession: B69679
A:Status: nucleic acid sequence not shown; translation not shown
A:Molecule type: DNA
A:Residues: 1-4427 <KUN>

A:Cross-references: GB:Z99113; GB:AL009126; NID:g2634090; PIDN:CAB13602.1; PID:g2634102
A:Experimental source: strain 168
C:Comment: This enzyme is composed of four synthase units. Unit1 comprises beta-ketosynt
acyl-carrier protein domains. Unit3 comprises beta-ketosynthase, acyl-carrier protein an
C:Genetics:
A:Gene: pksL; pksX
C:Superfamily: Bacillus subtilis polyketide synthase pksL; 3-oxoacyl-[acyl-carrier-prote
C:Keywords: acyltransferase; carrier protein
F:343-758/Domain: 3-oxoacyl-[acyl-carrier-protein] synthase I homology <OAS1>
F:1410-1591/Domain: short-chain alcohol dehydrogenase homology <SAD1>
F:1836-2252/Domain: 3-oxoacyl-[acyl-carrier-protein] synthase I homology <OAS2>
F:2485-2559/Domain: acyl carrier protein homology <ACP2>
F:2636-2700/Domain: acyl carrier protein homology <ACP2>
F:2783-3181/Domain: 3-oxoacyl-[acyl-carrier-protein] synthase I homology <OAS3>
F:3576-3774/Domain: short-chain alcohol dehydrogenase homology <SAD2>
F:3852-3922/Domain: acyl carrier protein homology <ACP3>
F:3992-4372/Domain: 3-oxoacyl-[acyl-carrier-protein] synthase I homology <OAS4>
Query Match 61.8%; Score 34; DB 2; Length 4427;
Best Local Similarity 100.0%; Pred. No. 6.6e+02;
Matches 0; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLV 6
|||||
DB 691 HHQKLV 696
RESULT 45
S39644
acetoin utilization protein acuD - Bacillus subtilis
C:Species: Bacillus subtilis
C:Date: 08-Jun-1994 #sequence_revision 10-Nov-1995 #text_change 21-Jul-2000
C:Accession: S39644; D69582
R:Grundv, F.J.; Waters, D.A.; Takova, T.Y.; Henkin, T.M.
Mol. Microbiol. 10, 259-271, 1993
A:Title: Identification of genes involved in utilization of acetate and acetoin in Bacil
A:Reference number: S39641; MUID:95020526
A:Accession: S39644
A>Status: preliminary
A:Molecule type: DNA
A:Residues: 1-214 <GR>
A:Cross-references: GB:L17309; NID:g861173; PIDN:AAA68285.1; PID:g348051
R:Kunst, F.; Ogasawara, N.; Moszer, I.; Albertini, A.M.; Alloni, G.; Azevedo, V.; Berte
C:Bron, S.; Brouillet, S.; Brusch, C.V.; Caldwell, B.; Capuano, V.; Carter, N.M.; Ch
A:Ehrlich, S.D.; Emerson, P.T.; Entian, K.D.; Errington, J.; Fabret, C.; Ferrari, E.
Nature 390, 249-256, 1997
A:Authors: Foulger, D.; Fritz, C.; Fujita, M.; Fujita, Y.; Fuma, S.; Galizzi, A.; Gall
Iech, J.; Harwood, C.R.; Henaut, A.; Hilbert, H.; Holsappel, S.; Hosono, S.; Hulio, M.F.
Koetter, P.; Koningsstein, G.; Krogh, M.; Kumano, M.; Kurita, K.; Lapidus, A.; Lardinols
A:Authors: Lauber, J.; Lazarevic, V.; Lee, S.M.; Levine, A.; Liu, H.; Masuda, S.; Maueel
Y, M.; Ogawa, K.; Ogiwara, A.; Oudega, B.; Park, S.H.; Parro, V.; Pohl, T.M.; Portetelle
Rieger, M.; Rivolta, C.; Rocha, E.; Roche, B.; Rose, M.; Sadale, Y.; Sato, T.; Scanlon
A:Authors: Schleich, S.; Schroeter, R.; Scoffone, F.; Sekiguchi, J.; Sekowska, A.; Seron
akeuchi, M.; Tamakoshi, A.; Tanaka, T.; Terpstra, P.; Tognoni, A.; Tosato, V.; Uchiyama,
T.; Winters, P.; Wipat, A.; Yamamoto, H.; Yamane, K.; Yasumoto, K.; Yata, K.; Yoshida, K
A:Authors: Yoshikawa, H.F.; Zumstein, E.; Yoshikawa, H.; Danchin, A.
A:Title: The complete genome sequence of the Gram-positive bacterium Bacillus subtilis.
A:Reference number: A69580; MUID:98044033
A:Accession: D69582
A>Status: preliminary; nucleic acid sequence not shown; translation not shown
A:Molecule type: DNA
A:Residues: 1-214 <KUN>
A:Cross-references: GB:Z99119; GB:AL009126; NID:g2635411; PIDN:CAB14948.1; PID:el185843;
A:Experimental source: strain 168
C:Genetics:
A:Gene: acuD
Query Match 60.0%; Score 33; DB 2; Length 214;
Best Local Similarity 83.3%; Pred. No. 43;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLV 6
|||||

Db 110 HHQKLI 115
RESULT 46
S41511
Brn-3a protein - mouse
C:Species: Mus musculus (house mouse)
C:Date: 25-Dec-1994 #sequence_revision 01-Sep-1995 #text_change 17-Mar-1999
C:Accession: S41511
R:Theil, T.; McLean-Hunter, S.; Zoernig, M.; Mosroey, T.
Nucleic Acids Res. 21, 5921-5929, 1993
A:Title: Mouse Brn-3 family of POU transcription factors: a new aminoterminal domain
A:Reference number: S41511; MUID:94119691
A:Accession: S41511
A>Status: preliminary
A:Molecule type: mRNA
A:Residues: 1-255 <THE>
C:Superfamily: unassigned homeobox proteins; homeobox homology; POU domain homology
C:Keywords: DNA binding; homeobox; nucleus; transcription regulation
F:110-180/Domain: POU domain homology <POU>
F:199-255/Domain: homeobox homology <HOX>
Query Match 60.0%; Score 33; DB 2; Length 255;
Best Local Similarity 60.0%; Pred. No. 52;
Matches 6; Conservative 2; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFAE 10
|||||
DB 107 HHRELEFAE 116
RESULT 47
D72217
conserved hypothetical protein - Thermotoga maritima (strain MSB8)
C:Species: Thermotoga maritima
C:Date: 11-Jun-1999 #sequence_revision 11-Jun-1999 #text_change 21-Jul-2000
C:Accession: D72217
R:Nelson, K.E.; Clayton, R.A.; Gill, S.R.; Gwinn, M.L.; Dodson, R.J.; Haft, D.H.; Hic
Garrett, M.M.; Stewart, A.M.; Cotton, M.D.; Pratt, M.S.; Phillips, C.A.; Richardson,
C.M.
Nature 399, 323-329, 1999
A:Title: Evidence for lateral gene transfer between Archaea and Bacteria from genome
A:Reference number: A72200; MUID:99287316
A:Accession: D72217
A>Status: preliminary
A:Molecule type: DNA
A:Residues: 1-258 <ARN>
A:Cross-references: GB:AE001812; GB:AE000512; NID:g4982302; PIDN:AAD36798.1; PID:g498
A:Experimental source: strain MSB8
C:Genetics:
A:Gene: TM1733
C:Superfamily: conserved hypothetical protein HI0072
Query Match 60.0%; Score 33; DB 2; Length 258;
Best Local Similarity 50.0%; Pred. No. 53;
Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;
QY 1 HHQKLVFAE 10
|||||
DB 140 HHSSMWFFAD 149
RESULT 48
A47003
cytokine receptor family class II protein CRF2-4 precursor - human
C:Species: Homo sapiens (man)
C:Date: 09-Sep-1994 #sequence_revision 09-Sep-1994 #text_change 01-Dec-2000
C:Accession: A47003; G01418
R:Lutfalla, G.; Gardner, K.; Uze, G.
Genomics 16, 366-373, 1993
A:Title: A new member of the cytokine receptor gene family maps on chromosome 21 at 1
A:Reference number: A47003; MUID:93300510
A:Accession: A47003

A:Status: preliminary
A:Molecule type: mRNA
A:Residues: 1-325 <LUT>
A:Cross-references: GB:Z17227; NID:g393378; PIDN:CAA78933.1; PID:g393379
R:Lutfalla, G.
submitted to the EMBL Data Library, April 1994

A:Reference number: G08935
A:Accession: G01418
A:Status: preliminary; translated from GB/EMBL/DDBJ
A:Molecule type: DNA
A:Residues: 1-123, 'D', '125-268, 'VGRME' <LU2>
A:Cross-references: EMBL:U08988; NID:g571295; PID:g571296

C:Genetics:
A:Gene: GDB:CRFB4; CRF2-4
A:Cross-references: GDB:138168; OMIM:123889
A:Map position: 21q; 21q22.1-21q22.2
A:Introns: 17/1; 58/2; 111/1; 166/3; 216/1
C:Keywords: transmembrane protein

Query Match 60.0%; Score 33; DB 2; Length 325;
Best Local Similarity 55.6%; Pred. No. 67;
Matches 5; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLYFFA 9
|| | | | |
Db 274 HHNTLLFFS 282

RESULT 49
T20562
hypothetical protein F07H5.2 - Caenorhabditis elegans
C:Species: Caenorhabditis elegans
C>Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 15-Oct-1999
C:Accession: T20562
R:Steward, C.
submitted to the EMBL Data Library, December 1995
A:Reference number: Z19292
A:Accession: T20562
A:Status: preliminary; translated from GB/EMBL/DDBJ
A:Molecule type: DNA
A:Residues: 1-334 <WIL>
A:Cross-references: EMBL:Z68314; PIDN:CAA92663.1; GSPDB:GN00020; CESP:F07H5.2
A:Experimental source: clone F07H5
C:Genetics:
A:Gene: CESP:F07H5.2
A:Map position: 2
A:Introns: 72/2; 146/3; 217/1; 280/3

Query Match 60.0%; Score 33; DB 2; Length 334;
Best Local Similarity 75.0%; Pred. No. 69;
Matches 6; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFFAE 10
||| | | | | |
Db 195 QKLFFAD 202

RESULT 50
S32170
phytoene synthetase - Myxococcus xanthus
C:Species: Myxococcus xanthus
C>Date: 06-Jan-1995 #sequence_revision 06-Jan-1995 #text_change 02-Mar-2001
C:Accession: S32170; S67951
R:Botella, J.; Murillo, F.; Ruiz-vazquez, R.
submitted to the EMBL Data Library, March 1993
A:Description: Nucleotide and deduced protein sequences of a carotenoid gene cluster in
A:Reference number: S32168
A:Accession: S32170
A:Molecule type: DNA
A:Residues: 1-336 <BOT>
A:Cross-references: EMBL:Z21955; NID:g577589; PIDN:CAA79957.1; PID:g288222
A:Experimental source: strain DK1050
R:Botella, J.A.; Murillo, F.J.; Ruiz-Vazquez, R.

Eur. J. Biochem. 233, 238-248, 1995
A:Title: A cluster of structural and regulatory genes for light-induced carotenogenes
A:Reference number: S67950; MUID:96061955
A:Accession: S67951
A:Molecule type: DNA
A:Residues: 151-175; 185-213 <BOW>
C:Genetics:
A:Start codon: GTG
C:Superfamily: Mycobacterium marinum phytoene synthase

Query Match 60.0%; Score 33; DB 2; Length 336;
Best Local Similarity 66.7%; Pred. No. 70;
Matches 6; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 1 HHQKLYFFA 9
|| | | | |
Db 23 HHAKSFFFA 31

Search completed: October 29, 2002, 09:24:30
Job time : 19 secs

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:23:27 ; Search time 11 seconds
(without alignments)
35.200 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 105224 seqs, 38719550 residues

Total number of hits satisfying chosen parameters: 105224

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : SwissProt_40:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Query	Score	Match	Length	DB	ID	Description
1	55	100.0	57	1	A4_PIG	Q29023	sus scrofa
2	55	100.0	57	1	A4_URSA	Q29149	ursus marit
3	55	100.0	58	1	A4_CANFA	Q28280	canis famil
4	55	100.0	58	1	A4_RABIT	Q28748	oryctolagus
5	55	100.0	58	1	A4_SHEEP	Q28757	ovis aries
6	55	100.0	59	1	A4_BOVIN	Q28053	bos taurus
7	55	100.0	751	1	A4_SAISC	Q95241	saimiri sci
8	55	100.0	770	1	A4_HUMAN	P05067	homo sapien
9	47	85.5	770	1	A4_MOUSE	P12023	mus musculus
10	47	85.5	770	1	A4_RAT	P08592	rattus norv
11	39	70.9	699	1	MALO_HAEIN	P45176	haemophilus
12	38	69.1	549	1	G6PI_ECOLI	P11537	escherichia
13	38	69.1	550	1	G6PI_VIBCH	Q9KUY4	vibrio chol
14	36	65.5	109	1	SAS_PIG	Q29257	sus scrofa
15	36	65.5	204	1	TNBE6_HUMAN	Q95857	homo sapien
16	36	65.5	210	1	SAS_HUMAN	P12999	homo sapien
17	35	63.6	549	1	G6PI_BUCAL	P57636	buchnera ap
18	35	63.6	549	1	G6PI_PASMU	Q9CN12	pasteurella
19	35	63.6	849	1	RS62_HUMAN	P15283	homo sapien
20	35	63.6	2347	1	KROS_HUMAN	P08922	homo sapien
21	34	61.8	124	1	SLP_BACSU	P39910	bacillus su
22	34	61.8	763	1	YN51_YEAST	P42843	saccharomyc
23	34	61.8	1163	1	RPOD_PEA	P12227	pisum sativ
24	34	61.8	1356	1	ROM2_YEAST	P51862	saccharomyc
25	34	61.8	2715	1	TRX2_HUMAN	Q9UMM6	homo sapien
26	34	61.8	4427	1	PKSL_BACSU	Q05470	bacillus su
27	33	60.0	214	1	ACUB_BACSU	P39066	bacillus su
28	33	60.0	258	1	PPNK_THEMA	Q9X255	thermotoga
29	33	60.0	325	1	ILOS_HUMAN	Q08334	homo sapien
30	33	60.0	496	1	G7D9_SOYBN	O81971	glycine max
31	33	60.0	549	1	G6PI_HAEIN	P44312	haemophilus
32	33	60.0	564	1	SVT_MYCGE	P47615	mycoplasma
33	33	60.0	2150	1	SDC3_CAEEEL	P34706	caenorhabd1

34	32	58.2	167	1	G6PI_KLEOX	P77877	klebsiella
35	32	58.2	178	1	CALC_MOUSE	O63811	mus musculus
36	32	58.2	321	1	CYF_GUTH	O78494	guillardia
37	32	58.2	380	1	FDSE_SOYBN	P48625	glycine max
38	32	58.2	481	1	LBP_HUMAN	P18428	homo sapien
39	32	58.2	492	1	CPBJ_MOUSE	O55071	mus musculus
40	32	58.2	495	1	MURC_RICPR	Q9ZDS8	rickettsia
41	32	58.2	501	1	ACHB_MOUSE	P09690	mus musculus
42	32	58.2	517	1	ACHG_HUMAN	P07510	homo sapien
43	32	58.2	519	1	ACHG_BOVIN	P13536	bos taurus
44	32	58.2	519	1	ACHG_MOUSE	P04760	mus musculus
45	32	58.2	519	1	ACHG_RAT	P18916	rattus norv
46	32	58.2	557	1	PRXV_ASCNO	P81701	ascophyllum
47	32	58.2	616	1	YANG_SCHPO	Q10190	schizosacch
48	32	58.2	734	1	GLGB_AGRTU	P52979	agrobacteri
49	32	58.2	808	1	PLD_PIMBR	O04883	pimpinella
50	32	58.2	808	1	PLD_RICCO	Q41142	ricinus com

ALIGNMENTS

RESULT 1

ID A4_PIG STANDARD; PRT; 57 AA.

AC Q29023;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Sus scrofa (Pig).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
OX NCBI_TaxID=9823;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Brain;
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid peptide in dog, polar bear and five other mammals by cross-species polymerase chain reaction analysis.";
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.

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CC EMBL; X56127; CAA39592.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
FT NON_TER 1
FT CHAIN 6 48 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN <1 33 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 34 57 POTENTIAL.
FT NON_TER 57 57
SQ SEQUENCE 57 AA; 6172 MW; 84209D88EBA82DFA CRC64;

Query Match 100.0%; Score 55; DB 1; Length 57;

Best Local Similarity 100.0%; Pred. No. 0.00019;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 18 HHQKLVFFAE 27

RESULT 2

A4_URSWA
ID A4_URSWA STANDARD; PRT; 57 AA.
AC Q29149; (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid
protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Ursus maritimus (Polar bear) (Thalarchos maritimus).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Carnivora; Fissipedia; Ursidae; Ursus.
OX NCBI_TaxID=29073;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Brain;
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid
peptide in dog, polar bear and five other mammals by cross-species
polymerase chain reaction analysis";
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -1- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
G(O) (BY SIMILARITY).
CC -1- SUBCELLULAR LOCATION: Type I membrane protein.
CC -1- SIMILARITY: BELONGS TO THE APP FAMILY.

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or send an email to license@isb-sib.ch).
CC
CC EMBL; X56125; CAA39593.1; -
CC HSP; P05067; IAML.
CC InterPro; IPR001868; A4_APP.
CC PROSITE; PS00319; A4_EXTRA; PARTIAL.
CC PROSITE; PS00320; A4_INTRA; PARTIAL.
CC Glycoprotein; Amyloid; Neurone; Transmembrane.
FT CHAIN 1 1
FT NON_TER 1 1
FT DOMAIN <1 33 BETA-AMYLOID PROTEIN (POTENTIAL).
FT TRANSMEM 34 57 EXTRACELLULAR (POTENTIAL).
FT NON_TER 57 57
FT SEQUENCE 57 AA; 6172 MW; 84209D88EA82DFA CRC64;
Query Match 100.0%; Score 55; DB 1; Length 57;
Best Local Similarity 100.0%; Pred. No. 0.00019;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 18 HHQKLVFFAE 27

RESULT 3

A4_CANFA
ID A4_CANFA STANDARD; PRT; 58 AA.
AC Q28280;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)

DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid
protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Canis familiaris (Dog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis.
OX NCBI_TaxID=9615;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Kidney;
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid
peptide in dog, polar bear and five other mammals by cross-species
polymerase chain reaction analysis";
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -1- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
G(O) (BY SIMILARITY).
CC -1- SUBCELLULAR LOCATION: Type I membrane protein.
CC -1- SIMILARITY: BELONGS TO THE APP FAMILY.

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CC
CC EMBL; X56125; CAA39590.1; -
CC HSP; P05067; IBA4.
CC InterPro; IPR001868; A4_APP.
CC PROSITE; PS00319; A4_EXTRA; PARTIAL.
CC PROSITE; PS00320; A4_INTRA; PARTIAL.
CC Glycoprotein; Amyloid; Neurone; Transmembrane.
FT CHAIN 1 1
FT NON_TER 1 1
FT DOMAIN <1 34 BETA-AMYLOID PROTEIN (POTENTIAL).
FT TRANSMEM 35 58 EXTRACELLULAR (POTENTIAL).
FT NON_TER 58 58
FT SEQUENCE 58 AA; 6285 MW; 8469D488A2E12DFA CRC64;
Query Match 100.0%; Score 55; DB 1; Length 58;
Best Local Similarity 100.0%; Pred. No. 0.0002;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 19 HHQKLVFFAE 28

RESULT 4

ID A4_RABIT STANDARD; PRT; 58 AA.
AC Q28748;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid
protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Oryctolagus cuniculus (Rabbit).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Lagomorpha; Leporidae; Oryctolagus.
OX NCBI_TaxID=9986;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Brain;
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid

QY 1 HHQKLVFFAE 10
DB 18 HHQKLVFFAE 27


```
RT peptide in dog, polar bear and five other mammals by cross-species
RT polymerase chain reaction analysis."
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
CC INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN
CC G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -----
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CC -----
DR EMBL; X56129; CAA39594.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
FT NON_TER 1
FT CHAIN 1 6 48 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN <1 33 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 34 57 POTENTIAL.
FT DOMAIN 58 >58 CYTOPLASMIC (POTENTIAL).
FT NON_TER 58 58
FT SEQUENCE 58 AA; 6300 MW; F434209D88EBA82D CRC64;
DR EMBL; X56129; CAA39594.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
FT NON_TER 1
FT CHAIN 1 6 48 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN <1 33 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 34 57 POTENTIAL.
FT DOMAIN 58 >58 CYTOPLASMIC (POTENTIAL).
FT NON_TER 58 58
FT SEQUENCE 58 AA; 6300 MW; F434209D88EBA82D CRC64;
Query Match 100.0%; Score 55; DB 1; Length 58;
Best Local Similarity 100.0%; Pred. No. 0.0002;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 18 HHQKLVFFAE 27
RESULT 5
A4_SHEEP
ID A4_SHEEP STANDARD; PRT; 58 AA.
AC Q28757;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid
DE protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Ovis aries (Sheep).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae;
OC Bovidae; Caprinae; Ovis.
OX NCBI_TaxID=9940;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Heart.
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid
RT peptide in dog, polar bear and five other mammals by cross-species
RT polymerase chain reaction analysis."
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
CC INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN
CC G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -----
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CC -----
DR EMBL; X56129; CAA39594.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
FT NON_TER 1
FT CHAIN 1 6 48 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN <1 33 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 34 57 POTENTIAL.
FT DOMAIN 58 >58 CYTOPLASMIC (POTENTIAL).
FT NON_TER 58 58
FT SEQUENCE 58 AA; 6300 MW; F434209D88EBA82D CRC64;
Query Match 100.0%; Score 55; DB 1; Length 58;
Best Local Similarity 100.0%; Pred. No. 0.0002;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 18 HHQKLVFFAE 27
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CC -----
DR EMBL; X56130; CAA39595.1; -.
DR HSSP; P05067; 1AHL.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
FT NON_TER 1
FT CHAIN 1 6 48 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN <1 33 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 34 57 POTENTIAL.
FT DOMAIN 58 >58 CYTOPLASMIC (POTENTIAL).
FT NON_TER 58 58
FT SEQUENCE 58 AA; 6300 MW; F434209D88EBA82D CRC64;
Query Match 100.0%; Score 55; DB 1; Length 58;
Best Local Similarity 100.0%; Pred. No. 0.0002;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 18 HHQKLVFFAE 27
RESULT 6
A4_BOVIN
ID A4_BOVIN STANDARD; PRT; 59 AA.
AC Q28053;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog [Contains: Beta-amyloid
DE protein (Beta-APP) (A-beta)] (Fragment).
GN APP.
OS Bos taurus (Bovine).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae;
OC Bovidae; Bovinae; Bos.
OX NCBI_TaxID=9913;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Brain.
RX MEDLINE=92017079; PubMed=1656157;
RA Johnstone E.M., Chaney M.O., Norris F.H., Pascual R., Little S.P.;
RT "Conservation of the sequence of the Alzheimer's disease amyloid
RT peptide in dog, polar bear and five other mammals by cross-species
RT polymerase chain reaction analysis."
RL Brain Res. Mol. Brain Res. 10:299-305(1991).
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
CC INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN
CC G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -----
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CC -----
DR EMBL; X56124; CAA39589.1; -.
DR EMBL; X56126; CAA39591.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR PROSITE; PS00319; A4_EXTRA; PARTIAL.
DR PROSITE; PS00320; A4_INTRA; PARTIAL.
KW Glycoprotein; Amyloid; Neurone; Transmembrane.
```

FT NON_TER 1 1
 FT CHAIN 7 49 BETA-AMYLOID PROTEIN (POTENTIAL).
 FT DOMAIN <1 34 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 35 58 POTENTIAL.
 FT DOMAIN 59 >59 CYTOPLASMIC (POTENTIAL).
 FT NON_TER 59 59
 SQ SEQUENCE 59 AA; 6414 MW; F43469D488A2E12D CRC64;

 Query Match 100.0%; Score 55; DB 1; Length 59;
 Best Local Similarity 100.0%; Pred. No. 0.0002;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

 QY 1 HHOKLVFFAE 10
 Db 19 HHOKLVFFAE 28
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 RESULT 7
 A4_SAISC STANDARD; PRT; 751 AA.
 AC Q95241;
 DT 15-DEC-1998 (Rel. 37, Created)
 DT 15-DEC-1998 (Rel. 37, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Alzheimer's disease amyloid A4 protein precursor [Contains: Beta-amyloid protein (Beta-APP) (A-beta)].
 GN APP.
 OS Saimiri sciureus (Common squirrel monkey).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Platyrrhini; Cebidae; Cebinae; Saimiri.
 OX NCBI_TaxID=9521;
 RN [1]
 RP SEQUENCE FROM N.A.
 RC TISSUE=Liver, and Kidney;
 RX MEDLINE=96108492; PubMed=8532114;
 RA Levy E., Amorim A., Frangione B., Walker L.C.;
 RT "Beta-amyloid precursor protein gene in squirrel monkeys with cerebral amyloid angiopathy.";
 RL Neurobiol. Aging 16:805-808(1995).
 CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN G(O).
 CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
 CC -!- DOMAIN: THE CLATHRIN-BINDING SITE IS ESSENTIAL FOR ITS ASSOCIATION WITH X11-ALPHA, -BETA, AND -GAMMA. THE SEQUENCE SPECIFIC RECOGNITION EXTENDS TO PEPTIDE RESIDUES THAT ARE C-TERMINAL TO THE NPXY MOTIF. THIS INTERACTION APPEARS TO BE INDEPENDENT OF PHOSPHORYLATION (BY SIMILARITY).
 CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
 CC -!- SIMILARITY: CONTAINS 1 BPTI/KUNITZ INHIBITOR DOMAIN.

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 CC EMBL; S81024; AAD14347.1; -
 CC HSSP; P05067; 1AAP.
 DR InterPro; IPR001868; A4_APP.
 DR InterPro; IPR002223; Kunitz_BPTI.
 DR Pfam; PF021177; A4_EXTRA; 1.
 DR Pfam; PF00014; Kunitz_BPTI; 1.
 DR PRINTS; PR00203; AMYLOIDA4.
 DR PRINTS; PR00759; BASICPTASE.
 DR SMART; SM00006; A4_EXTRA; 1.
 DR SMART; SM00131; KU; 1.
 DR PROSITE; PS00319; A4_EXTRA; 1.
 DR PROSITE; PS00320; A4_INTRA; 1.
 DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
 DR PROSITE; PS00279; BPTI_KUNITZ_2; 1.

KW Glycoprotein; Amyloid; Neurone; Transmembrane; Alternative splicing;
 KW Signal; Serine protease inhibitor.
 FT SIGNAL 1 17 BY SIMILARITY.
 FT CHAIN 18 751 A4 PROTEIN.
 FT CHAIN 653 695 BETA-AMYLOID PROTEIN (POTENTIAL).
 FT DOMAIN 18 680 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 681 704 POTENTIAL.
 FT DOMAIN 705 751 CYTOPLASMIC (POTENTIAL).
 FT DOMAIN 287 345 BPTI/KUNITZ INHIBITOR.
 FT SITE 740 743 CLATHRIN-BINDING (BY SIMILARITY).
 FT ACT_SITE 301 302 REACTIVE BOND.
 FT DISULFID 291 341 BY SIMILARITY.
 FT DISULFID 300 324 BY SIMILARITY.
 FT DISULFID 316 337 BY SIMILARITY.
 FT CARBOHYD 523 523 N-LINKED (GLCNAC. . .) (PROBABLE).
 FT CARBOHYD 552 552 N-LINKED (GLCNAC. . .) (PROBABLE).
 SQ SEQUENCE 751 AA; 84893 MW; 6C3E431089569049 CRC64;

 Query Match 100.0%; Score 55; DB 1; Length 751;
 Best Local Similarity 100.0%; Pred. No. 0.0027;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

 QY 1 HHOKLVFFAE 10
 Db 665 HHOKLVFFAE 674
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 RESULT 8
 A4_HUMAN STANDARD; PRT; 770 AA.
 AC P05067; P09000; Q16011;
 DT 13-AUG-1987 (Rel. 05, Created)
 DT 01-NOV-1991 (Rel. 20, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Alzheimer's disease amyloid A4 protein precursor (Protease nexin-II) (PN-II) (APPI) [Contains: Beta-amyloid protein (Beta-APP) (A-beta)].
 DE APP OR A4 OR CVAP OR AD1.
 GN Homo sapiens (Human).
 OS Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
 OX NCBI_TaxID=9606;
 RN [1]
 RP SEQUENCE FROM N.A.
 RC TISSUE=Brain;
 RX MEDLINE=87144572; PubMed=2881207;
 RA Kang J., Lemaire H.-G., Unterbeck A., Salbaum J.M., Masters C.L., Grzeschik K.-H., Multhaup G., Beyreuther K., Mueller-Hill B.;
 RT "The precursor of Alzheimer's disease amyloid A4 protein resembles a cell-surface receptor.";
 RT Nature 325:733-736(1987).
 RL [2]
 RN SEQUENCE FROM N.A.
 RP MEDLINE=88122639; PubMed=2893289;
 RX Ponte P., Gonzalez-Dewhitt P., Schilling J., Miller J., Hsu D., Greenberg B., Davis K., Wallace W., Lieberburg I., Fuller F., Cordell B.;
 RA "A new A4 amyloid mRNA contains a domain homologous to serine protease inhibitors.";
 RT Nature 331:525-527(1988).
 RL [3]
 RN SEQUENCE FROM N.A.
 RP MEDLINE=89128427; PubMed=2783775;
 RX Lemaire H.G., Salbaum J.M., Multhaup G., Kang J., Bayne R.M., Unterbeck A., Beyreuther K., Mueller-Hill B.;
 RA "The preA4(695) precursor protein of Alzheimer's disease A4 amyloid is encoded by 16 exons.";
 RT Nucleic Acids Res. 17:517-522(1989).
 RL [4]
 RN SEQUENCE FROM N.A.
 RP MEDLINE=97263807; PubMed=9108164;
 RX Hattori M., Tsukahara F., Furuhashi Y., Tanahashi H., Hirose M., Saito M., Tsukuni S., Sakaki Y.;
 RA "A novel method for making nested deletions and its application for

RT sequencing of a 300 kb region of human APP locus.";
RL Nucleic Acids Res. 25:1802-1808(1997).
RP SEQUENCE OF 286-345 AND 365-366 FROM N.A.
RX MEDLINE-8812640; PubMed-2893290;
RA Tanzi R.E., McClatchey A.I., Lamperti E.D., Villa-Komaroff L.,
RA Gusella J.F., Neve R.L.;
RT "Protease inhibitor domain encoded by an amyloid protein precursor
RT mRNA associated with Alzheimer's disease.";
RL Nature 331:528-530(1988).
RN [6]
RP SEQUENCE OF 287-367 FROM N.A.
RX MEDLINE-8812641; PubMed-2893291;
RA Kitaguchi N., Takahashi Y., Tokushima Y., Shiojiri S., Ito H.;
RT "Novel precursor of Alzheimer's disease amyloid protein shows
RT protease inhibitory activity.";
RL Nature 331:530-532(1988).
RN [7]
RP SEQUENCE OF 284-289 AND 365-770 FROM N.A.
RX MEDLINE-87231971; PubMed-3035574;
RA Robakis N.K., Ramakrishna N., Wolfe G., Wisniewski H.M.;
RT "Molecular cloning and characterization of a cDNA encoding the
RT cerebrovascular and the neuritic plaque amyloid peptides.";
RL Proc. Natl. Acad. Sci. U.S.A. 84:4190-4194(1987).
RN [8]
RP SEQUENCE OF 507-770 FROM N.A.
RX MEDLINE-88124954; PubMed-2893379;
RA Zain S.B., Salim M., Chou W.G., Sajdel-Sulkowska E.M., Majocha R.E.,
RA Marotta C.A.;
RT "Molecular cloning of amyloid cDNA derived from mRNA of the Alzheimer
RT disease brain: coding and noncoding regions of the fetal precursor
RT mRNA are expressed in the cortex.";
RL Proc. Natl. Acad. Sci. U.S.A. 85:929-933(1988).
RN [9]
RP SEQUENCE OF 672-681.
RX MEDLINE-88035004; PubMed-3312495;
RA Partridge W.M., Vinters H.V., Yang J., Eisenberg J., Choi T.B.,
RA Tourtellotte W.W., Huebner V., Shively J.E.;
RT "Amyloid angiopathy of Alzheimer's disease: amino acid composition
RT and partial sequence of a 4,200-dalton peptide isolated from cortical
RT microvessels.";
RL J. Neurochem. 49:1394-1401(1987).
RN [10]
RP SEQUENCE OF 739-770 FROM N.A.
RX MEDLINE-90236318; PubMed-2110105;
RA Yoshikai S.-I., Sasaki H., Doh-Ura K., Furuya H., Sakaki Y.;
RT "Genomic organization of the human amyloid beta-protein precursor
RT gene.";
RL Gene 87:257-263(1990).
RN [11]
RP SEQUENCE OF 1-10 FROM N.A.
RX TISSUE-Liver;
RA Schon E.A., Mita S., Sadlock J., Herbert J.;
RT "A cDNA specifying the human amyloid beta precursor protein (ABPP)
RT encodes a 95-kDa polypeptide.";
RL Nucleic Acids Res. 16:9351-9351(1988).
RN [12]
RP SEQUENCE OF 18-50.
RX MEDLINE-87250462; PubMed-3597385;
RA van Nostrand W.E., Cunningham D.D.;
RT "Purification of protease nexin II from human fibroblasts.";
RL J. Biol. Chem. 262:8508-8514(1987).
RN [13]
RP IDENTITY OF APP WITH NEXIN-II.
RX MEDLINE-89384866; PubMed-2506449;
RA Oltersdorf T., Fritz L.C., Schenk D.B., Lieberburg I.,
RA Johnson-Wood K.L., Beattie E.C., Ward P.J., Blacher R.W., Dovey H.F.,
RA Sinha S.;
RT "The secreted form of the Alzheimer's amyloid precursor protein with
RT the Kunitz domain is protease nexin-II.";
RL Nature 341:144-147(1989).
RN [14]

RP PROTEASE-SPECIFICITY OF INHIBITOR DOMAIN.
RX MEDLINE-90211252; PubMed-1969731;
RA Kido H., Fukutomi A., Schilling J., Wang Y., Cordell B., Katunuma N.;
RT "Protease-specificity of Kunitz inhibitor domain of Alzheimer's
RL disease amyloid protein precursor.";
RL Biochem. Biophys. Res. Commun. 167:716-721(1990).
RN [15]
RP COMPLEX WITH G(O).
RX MEDLINE-9318955; PubMed-8446172;
RA Nishimoto I., Okamoto T., Matsuura Y., Takahashi S., Okamoto T.,
RA Murayama Y., Ogata E.;
RT "Alzheimer amyloid protein precursor complexes with brain GTP-binding
RL protein G(O).";
RL Nature 362:75-79(1993).
RN [16]
RP X-RAY CRYSTALLOGRAPHY (1.8 ANGSTROMS) OF 28-133.
RX MEDLINE-99215582; PubMed-10201399;
RA Rossjohn J., Cappai R., Feil S.C., Henry A., McKinstry W.J.,
RA Galatis D., Hesse L., Multhaup G., Beyreuther K., Masters C.L.,
RA Parker M.W.;
RT "Crystal structure of the N-terminal, growth factor-like domain of
RT Alzheimer amyloid precursor protein.";
RL Nat. Struct. Biol. 6:327-331(1999).
RN [17]
RP X-RAY CRYSTALLOGRAPHY (1.5 ANGSTROMS) OF 287-344.
RX MEDLINE-91104913; PubMed-2125487;
RA Hynes T.R., Randal M., Kennedy L.A., Eigenbrot C., Kossiakof A.A.;
RT "X-ray crystal structure of the protease inhibitor domain of
RL Alzheimer's amyloid beta-protein precursor.";
RL Biochemistry 29:10018-10022(1990).
RN [18]
RP STRUCTURE BY NMR OF 289-344.
RX MEDLINE-92031488; PubMed-1718421;
RA Heald S.L., Tilton R.F. Jr., Hammond L.S., Lee A., Bayney R.M.,
RA Kamark M.E., Ramabhadran T.V., Dreyer R.N., Davis G., Unterbeck A.,
RA Tamburini P.P.;
RT "Sequential NMR resonance assignment and structure determination of
RT the Kunitz-type inhibitor domain of the Alzheimer's beta-amyloid
RL precursor protein.";
RL Biochemistry 30:10467-10478(1991).
RN [19]
RP STRUCTURE BY NMR OF 672-699.
RX MEDLINE-94281210; PubMed-7516706;
RA Talafous J., Marcinkowski K.J., Klopman G., Zagorski M.G.;
RT "Solution structure of residues 1-28 of the amyloid beta-peptide.";
RL Biochemistry 33:7788-7796(1994).
RN [20]
RP STRUCTURE BY NMR OF 696-706.
RX MEDLINE-97128622; PubMed-8973180;
RA Kohno T., Kobayashi K., Maeda T., Sato K., Takashima A.;
RT "Three-dimensional structures of the amyloid beta peptide (25-35) in
RL membrane-mimicking environment.";
RL Biochemistry 35:16094-16104(1996).
RN [21]
RP STRUCTURE BY NMR OF 672-711.
RX MEDLINE-98359783; PubMed-9693002;
RA Coles M., Bicknell W., Watson A.A., Fairlie D.P., Craik D.J.;
RT "Solution structure of amyloid beta-peptide(1-40) in a water-micelle
RL environment. Is the membrane-spanning domain where we think it is?";
RL Biochemistry 37:11064-11077(1998).
RN [22]
RP STRUCTURE BY NMR OF 672-699.
RX MEDLINE-20400066; PubMed-10940222;
RA Poulsen S.-A., Watson A.A., Craik D.J.;
RT "Solution structures in aqueous SDS micelles of two amyloid beta
RL peptides of Abeta(1-28) mutated at the alpha-secretase cleavage
RL site.";
RL J. Struct. Biol. 130:142-152(2000).
RN [23]
RP STRUCTURE BY NMR OF 681-706.
RX MEDLINE-20400065; PubMed-10940221;
RA Zhang S., Iwata K., Lachenmann M.J., Peng J.W., Li S., Stimson E.R.,
RA Lu Y., Felix A.M., Maggio J.E., Lee J.P.;

RT "The Alzheimer's peptide a beta adopts a collapsed coil structure in water.";
RT J. Struct. Biol. 130:130-141(2000).
RN [24]
RX SIGNAL SEQUENCE CLEAVAGE SITE, AND TOPOLOGY.
RX MEDLINE=89296437; PubMed=2900137;
RA Dykx T., Weidemann A., Multhaup G., Salbaum J.M., Lemaire H.-G.,
RA Kang J., Mueller-Hill B., Masters C.L., Beyreuther K.;
RT "Identification, transmembrane orientation and biogenesis of the amyloid A4 precursor of Alzheimer's disease.";
RT amyloid A4 precursor of Alzheimer's disease.";
Query Match 100.0%; Score 55; DB 1; Length 770;
Best Local Similarity 100.0%; Pred. No. 0.0027;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHOKLVFFAE 10
Db 684 HHOKLVFFAE 693
RESULT 9
A4_MOUSE STANDARD; PRT; 770 AA.
AC Fl2023;
DT 01-OCT-1989 (Rel. 12, Created)
DT 01-DEC-1992 (Rel. 24, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog precursor
DE (Amyloidogenic glycoprotein) (AG).
GN APP.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RN SEQUENCE OF 1-289 AND 365-770 FROM N.A.
RC STRAIN=BALE/C; TISSUE=Brain;
RX MEDLINE=92096458; PubMed=1756177;
RA de Strooper B., van Leuven F., van den Berghe H.;
RT "The amyloid beta protein precursor or proteinase nexin II from mouse is closer related to its human homolog than previously reported.";
RL Blochim. Biophys. Acta 1129:141-143(1991).
RN [2]
RN SEQUENCE OF 1-289 AND 365-770 FROM N.A.
RC TISSUE=Brain;
RX MEDLINE=88106489; PubMed=3322280;
RA Yamada T., Sasaki H., Furuya H., Miyata T., Goto I., Sakaki Y.;
RT "Complementary DNA for the mouse homolog of the human amyloid beta protein precursor.";
RL Biochem. Biophys. Res. Commun. 149:665-671(1987).
RN [3]
RN REVISIONS.
RA Yamada T.;
RL Submitted (MAR-1988) to the EMBL/GenBank/DBJ databases.
RN [4]
RN SEQUENCE OF 289-364 FROM N.A.
RC STRAIN=CD-1; TISSUE=Placenta;
RX MEDLINE=89345111; PubMed=2569710;
RA Fukuchi K., Martin G.M., Deeb S.S.;
RT "Sequence of the protease inhibitor domain of the A4 amyloid protein precursor of Mus domesticus.";
RL Nucleic Acids Res. 17:5396-5396(1989).
RN [5]
RN SEQUENCE OF 1-19 FROM N.A.
RX MEDLINE=92209998; PubMed=1555768;
RA Izumi R., Yamada T., Yoshikai S.I., Sasaki H., Hattori M., Sakai Y.;
RT "Positive and negative regulatory elements for the expression of the Alzheimer's disease amyloid precursor-encoding gene in mouse.";
RL Gene 112:189-195(1992).
RN [6]
RN SEQUENCE OF 281-380 FROM N.A., AND ALTERNATIVE SPLICING.
RC TISSUE=Brain, and Kidney;

RX MEDLINE=89149813; PubMed=2493250;
RA Yamada T., Sasaki H., Dohura K., Goto I., Sakaki Y.;
RT "Structure and expression of the alternatively-spliced forms of mRNA for the mouse homolog of Alzheimer's disease amyloid beta protein precursor.";
RL Biochem. Biophys. Res. Commun. 158:906-912(1989).
CC -1- SUBCELLULAR LOCATION: Type I membrane protein.
CC -1- ALTERNATIVE PRODUCTS: 5 ISOFORMS; APP(395), APP(563), APP(695), APP(751) AND APP(770) (SHOWN HERE); ARE PRODUCED BY ALTERNATIVE SPLICING.
CC -1- TISSUE SPECIFICITY: AAA(770) IS EXPRESSED IN KIDNEY. AAA(751) IS WIDELY EXPRESSED. AAA(695) IS EXPRESSED IN BRAIN, KIDNEY AND LIVER.
CC -1- DOMAIN: THE CLATHRIN-BINDING SITE IS ESSENTIAL FOR ITS ASSOCIATION WITH X11-ALPHA, -BETA, AND -GAMMA. THE SEQUENCE SPECIFIC RECOGNITION EXTENDS TO PEPTIDE RESIDUES THAT ARE C-TERMINAL TO THE NPXY MOTIF. THIS INTERACTION APPEARS TO BE INDEPENDENT OF PHOSPHORYLATION (BY SIMILARITY).
CC -1- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -1- SIMILARITY: CONTAINS 1 BPTI/KUNITZ INHIBITOR DOMAIN.
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CC
DR EMBL: X59379; -; NOT ANNOTATED_CDS.
DR EMBL: M18373; AAA37139.1; -;
DR EMBL: X15210; CAA33280.1; -;
DR EMBL: D10603; BAA01456.1; -;
DR EMBL: M24397; AAA39929.1; -;
DR PIR: A27485; A27485.
DR PIR: S04855; S04855.
DR PIR: S19727; S19727.
DR HSP: P05067; IQCM.
DR MGD: MGI:88059; App.
DR InterPro: IPR001868; A4_APP.
DR InterPro: IPR002223; Kunitz_BPTI.
DR Pfam: PF02177; A4_EXTRA; 1.
DR Pfam: PF00014; Kunitz_BPTI; 1.
DR PRINTS: PR00203; AMYLOIDAM.
DR PRINTS: PR00759; BASICPTASE.
DR SMART: SM00006; A4_EXTRA; 1.
DR SMART: SM00131; KU; 1.
DR PROSITE: PS00319; A4_EXTRA; 1.
DR PROSITE: PS00320; A4_INTRA; 1.
DR PROSITE: PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE: PS00279; BPTI_KUNITZ_2; 1.
KW Glycoprotein; Amyloid; Neurone; Transmembrane; Signal;
KW Alternative splicing; Serine protease inhibitor.
FT SIGNAL 1 17
FT CHAIN 18 770
FT ALZHEIMER'S DISEASE AMYLOID A4 PROTEIN
FT HOMOLOG.
FT DOMAIN 18 699
FT EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 700 723
FT POTENTIAL.
FT DOMAIN 724 770
FT CYTOPLASMIC (POTENTIAL).
FT DOMAIN 673 715
FT EQUIVALENT OF BETA-AMYLOID PROTEIN.
FT DOMAIN 287 345
FT BPTI/KUNITZ INHIBITOR.
FT SITE 759 762
FT CLATHRIN-BINDING (BY SIMILARITY).
FT DISULFID 291 341
FT BY SIMILARITY.
FT DISULFID 300 324
FT BY SIMILARITY.
FT DISULFID 316 337
FT BY SIMILARITY.
FT CARBOHYD 542 542
FT N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 571 571
FT N-LINKED (GLCNAC. . .) (POTENTIAL).
FT VARSPLIC 289 289
FT E -> V (IN ISOFORM APP(695)).
FT VARSPLIC 290 364
FT MISSING (IN ISOFORM APP(695)).
FT VARSPLIC 346 380
FT MISSING (IN ISOFORM APP(751)).
SQ SEQUENCE 770 AA; 86752 MW; 26C50DE0890CA7A CRC64;
Query Match 85.5%; Score 47; DB 1; Length 770;

Best Local Similarity 100.0%; Pred. No. 0.11;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFAE 10
Db 685 HOKLVFFAE 693
|||||

RESULT 10
ID A4_RAT STANDARD; PRT; 770 AA.
AC P08592;
DT 01-AUG-1988 (Rel. 08, Created)
DT 01-DEC-1992 (Rel. 24, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Alzheimer's disease amyloid A4 protein homolog precursor
DE (Amyloidogenic glycoprotein) (AG).
GN APP.
OS Rattus norvegicus (Rat).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX NCBI_TaxID=10116;
RN [1]
RP SEQUENCE OF 1-289 AND 365-770 FROM N.A.
RC TISSUE=Brain;
RX MEDLINE=88312583; PubMed=2900758;
RA Shivers B.D., Hilbich C., Multhaup G., Salbaum J.M., Beyreuther K.,
RA Seeburg P.H.;
RT "Alzheimer's disease amyloidogenic glycoprotein: expression pattern
in rat brain suggests a role in cell contact.";
RL EMBO J. 7:1365-1370(1988).
RN [2]
RP SEQUENCE OF 289-364 FROM N.A.
RC TISSUE=Liver;
RX MEDLINE=89183625; PubMed=2648331;
RA Kang J., Mueller-Hill B.;
RT "The sequence of the two extra exons in rat preA4.";
RL Nucleic Acids Res. 17:2130-2130(1989).
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- ALTERNATIVE PRODUCTS: 5 ISOFORMS; APP(395), APP(563), APP(695),
APP(751) AND APP(770) (SHOWN HERE); ARE PRODUCED BY ALTERNATIVE
SPlicing.
CC -!- DOMAIN: THE CLATHRIN-BINDING SITE IS ESSENTIAL FOR ITS ASSOCIATION
WITH X11-ALPHA, -BETA, AND -GAMMA. THE SEQUENCE SPECIFIC
RECOGNITION EXTENDS TO PEPTIDE RESIDUES THAT ARE C-TERMINAL TO THE
NPXY MOTIF. THIS INTERACTION APPEARS TO BE INDEPENDENT OF
PHOSPHORYLATION (BY SIMILARITY).
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -!- SIMILARITY: CONTAINS 1 BPTI/KUNITZ INHIBITOR DOMAIN.

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DR EMBL; X07648; CAA30488.1; -;
DR EMBL; X14066; CAA32229.1; -;
DR PIR; S00550; S00550.
DR PIR; S03607; S03607.
DR HSP; P05067; 1AAP.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOID4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00006; A4_EXTRA; 1.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.

DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS00279; BPTI_KUNITZ_2; 1.
KW Glycoprotein; Amyloid; Neurone; Transmembrane; Signal;
KW Alternative splicing; Serine protease inhibitor.
FT SIGNAL 1 17
FT CHAIN 18 770
FT FT
FT ALZHEIMER'S DISEASE AMYLOID A4 PROTEIN
FT HOMOLOG.
FT DOMAIN 18 699
FT EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 700 723
FT DOMAIN 724 770
FT POTENTIAL.
FT DOMAIN 673 715
FT CYTOPLASMIC (POTENTIAL).
FT DOMAIN 287 345
FT EQUIVALENT OF BETA-AMYLOID PROTEIN.
FT SITE 759 762
FT BPTI/KUNITZ INHIBITOR.
FT DISULFID 291 341
FT CLATHRIN-BINDING (BY SIMILARITY).
FT DISULFID 300 324
FT BY SIMILARITY.
FT DISULFID 316 337
FT BY SIMILARITY.
FT CARBOHYD 542 542
FT N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 571 571
FT N-LINKED (GLCNAC. . .) (POTENTIAL).
FT VARSPLIC 289 289
FT E -> V (IN ISOFORM APP(695)).
FT VARSPLIC 290 364
FT MISSING (IN ISOFORM APP(695)).
SQ SEQUENCE 770 AA; 86704 MW; C26C9D6B2D929A7 CRC64;
Query Match 85.5%; Score 47; DB 1; Length 770;
Best Local Similarity 100.0%; Pred. No. 0.11;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFAE 10
Db 685 HOKLVFFAE 693
|||||

RESULT 11
MALQ_HAEIN STANDARD; PRT; 699 AA.
ID MALQ_HAEIN
AC P45176;
DT 01-NOV-1995 (Rel. 32, Created)
DT 01-NOV-1995 (Rel. 32, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE 4-alpha-glucanotransferase (EC 2.4.1.25) (Amylomaltase)
DE (Disproportionating enzyme) (D-enzyme).
GN MALQ OR H11356.
OS Haemophilus influenzae.
OC Bacteria; Proteobacteria; gamma subdivision; Pasteurellaceae;
OC Haemophilus.
OX NCBI_TaxID=727;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=RD / KW20 / ATCC 51907;
RX MEDLINE=95350630; PubMed=7542800;
RA Fleischmann R.D., Adams M.D., White O., Clayton R.A., Kirkness E.F.,
RA Kerlavage A.R., Bult C.J., Tomb J.-F., Dougherty B.A., Merrick J.M.,
RA McKenney K., Sutton G., Fitzhugh W., Fields C.A., Gocayne J.D.,
RA Scott J.D., Shirley R., Liu L.-I., Glodek A., Keiley J.M.,
RA Weidman J.F., Phillips C.A., Spriggs T., Hedblom E., Cotton M.D.,
RA Utterback T.R., Hanna M.C., Nguyen D.T., Saudek D.M., Brandon R.C.,
RA Fine L.D., Fritchman J.L., Fuhrmann J.L., Geoghagen N.S.M.,
RA Gnehm C.L., McDonald L.A., Small K.V., Fraser C.M., Smith H.O.,
RA Venter J.C.;
RT "Whole-genome random sequencing and assembly of Haemophilus
influenzae Rd.";
RL Science 269:496-512(1995).
CC -!- CATALYTIC ACTIVITY: Transfers a segment of a (1,4)-alpha-D-glucan
to a new 4'-position in an acceptor, which may be glucose or (1,4)-
alpha-D-glucan.
CC -!- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
CC -!- SIMILARITY: BELONGS TO THE DISPROPORTIONATING ENZYME FAMILY.

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CC -----
DR EMBL; U32815; AAC23003.1; -.
DR TIGR; H11356; -.
DR InterPro; IPR003385; 4A_gluconotrans.
DR PFam; PF02446; 4A_gluconotrans; 1.
DR Transferrase; Glycosyltransferase; Carbohydrate metabolism;
KW Complete proteome.
SQ SEQUENCE 699 AA; 80251 MW; 80D6E1D51EC2E1E9 CRC64;

Query Match 70.9%; Score 39; DB 1; Length 699;
Best Local Similarity 66.7%; Pred. No. 3.8;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFFA 9
Db 349 HHEKIQFFA 357

RESULT 12
G6PI_ECOLI
ID G6PI_ECOLI STANDARD; PRT; 549 AA.
AC P11537;
DT 01-OCT-1989 (Rel. 12, Created)
DT 01-MAR-2002 (Rel. 41, Last sequence update)
DE Glucose-6-phosphate isomerase (EC 5.3.1.9) (GPI) (Phosphoglucose
DE isomerase) (PGI) (Phosphohexose isomerase) (PHI).
GN PGI OR B4025 OR Z5623 OR ECS5008.
OS Escherichia coli, and
OS Escherichia coli O157:H7.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562, 83334;
[1]
RN SEQUENCE FROM N.A.
RC STRAIN-JM101;
RX MEDLINE=89364675; PubMed=2549364;
RA Froman B.E., Tait R.C., Gottlieb L.D.;
RT "Isolation and characterization of the phosphoglucose isomerase gene
RT from Escherichia coli.";
RL Mol. Gen. Genet. 217:126-131(1989).
RN [2]
RP SEQUENCE FROM N.A., AND PHYLOGENETIC STUDY.
RC STRAIN-XLI BLUE 2;
RX MEDLINE=92277670; PubMed=1593646;
RA Smith M.W., Doolittle R.F.;
RT "Anomalous phylogeny involving the enzyme glucose-6-phosphate
RT isomerase.";
RL J. Mol. Evol. 34:544-545(1992).
RN [3]
RP SEQUENCE FROM N.A.
RC STRAIN-K12 / MG1655;
RX MEDLINE=94089392; PubMed=8265357;
RA Blattner F.R., Burland V.D., Plunkett G. III, Sofia H.J.,
RA Daniels D.L.;
RT "Analysis of the Escherichia coli genome. IV. DNA sequence of the
RT region from 89.2 to 92.8 minutes.";
RL Nucleic Acids Res. 21:5408-5417(1993).
RN [4]
RP SEQUENCE FROM N.A.
RC STRAIN-O157:H7 / EDL933 / ATCC 700927;
RX MEDLINE=21074935; PubMed=11206551;
RA Perina N.T., Plunkett G. III, Burland V., Mau B., Glasner J.D.,
RA Rose D.J., Mayhew G.F., Evans P.S., Gregor J., Kirkpatrick H.A.,
RA Posfai G., Hackett J., Klink S., Boutin A., Shao Y., Miller L.,
RA Grobeck E.J., Davis N.W., Lim A., Dimalanta E.T., Potamoudis K.,
RA Apodaca J., Anantharaman T.S., Lin J., Yen J., Schwartz D.C.,
RA Weich R.A., Blattner F.R.;
RT "Genome sequence of enterohaemorrhagic Escherichia coli O157:H7.";
RL Nature 409:529-533(2001).
RN [5]
RP SEQUENCE FROM N.A.
RC STRAIN-O157:H7 / RIMD 0509952;
RX MEDLINE=21156231; PubMed=11258796;
RA Hayashi T., Makino K., Ohnishi M., Kurokawa K., Ishii K., Yokoyama K.,
RA Han C.-G., Ohtsubo E., Nakayama K., Murata T., Tanaka M., Tobe T.,
RA Iida T., Takami H., Honda T., Sasakawa C., Ogasawara N., Yasunaga T.,
RA Kuhara S., Shiba T., Hattori M., Shinagawa H.;
RT "Complete genome sequence of enterohaemorrhagic Escherichia coli
RT O157:H7 and genomic comparison with a laboratory strain K-12.";
RL DNA Res. 8:11-22(2001).
CC -!- CATALYTIC ACTIVITY: D-glucose 6-phosphate = D-fructose 6-
CC phosphate.
CC -!- PATHWAY: INVOLVED IN GLYCOLYSIS AND IN GLUCONEOGENESIS.
CC -!- SUBUNIT: HOMODIMER.
CC -!- SUBCELLULAR LOCATION: Cytoplasmic.
CC -!- SIMILARITY: BELONGS TO THE GPI FAMILY.
CC -----
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CC or send an email to license@isb-sib.ch).
CC -----
CC EMBL; X15196; CAA33268.1; -.
CC EMBL; U00006; AAC43119.1; -.
CC DR EMBL; AE000476; AAC76995.1; -.
CC DR EMBL; AE005635; AAG59224.1; -.
CC DR EMBL; AP002568; BAB38431.1; -.
CC DR PIR; JS0142; NUC.
CC DR EcoGene; EGI0702; pgi.
CC DR InterPro; IPR001672; G6P_Isomerase.
CC DR Pfam; PF00342; PGI; 1.
CC DR PRINTS; PR00662; G6PISOMERASE.
CC DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
CC DR PROSITE; PS00174; P_GLUCOSE_ISOMERASE_2; 1.
CC KW isomerase; Gluconeogenesis; Glycolysis; Complete proteome.
FT ACT_SITE 386 386 BY SIMILARITY.
FT ACT_SITE 514 514 BY SIMILARITY.
FT CONFLICT 317 317 L -> V (IN REF. 1 AND 2).
SQ SEQUENCE 549 AA; 61529 MW; 74AEDB70A068A01 CRC64;

Query Match 69.1%; Score 38; DB 1; Length 549;
Best Local Similarity 66.7%; Pred. No. 4.6;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV-FFAE 10
Db 416 HHQKLLSNFFAQ 427

RESULT 13
G6PI_VIBCH
ID G6PI_VIBCH STANDARD; PRT; 550 AA.
AC Q9KUY4;
DT 01-MAR-2002 (Rel. 41, Created)
DT 01-MAR-2002 (Rel. 41, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Glucose-6-phosphate isomerase (EC 5.3.1.9) (GPI) (Phosphoglucose
DE isomerase) (PGI) (Phosphohexose isomerase) (PHI).
GN PGI OR VC0374.
OS Vibrio cholerae.
OC Bacteria; Proteobacteria; gamma subdivision; Vibrionaceae; Vibrio.
OX NCBI_TaxID=666;
[1]
RN SEQUENCE FROM N.A.
RC STRAIN-EL TOR N16961 / SEROTYPE O1;
RX MEDLINE=20406833; PubMed=10952301;
RA Heidelberg J.F., Eisen J.A., Nelson W.C., Clayton R.A., Ginn M.L.,
RA Dodson R.J., Haft D.H., Hickey E.K., Peterson J.D., Umayam L.A.,
RA Gill S.R., Nelson K.E., Read T.D., Tettelin H., Richardson D.,
RA Ermolaeva M.D., Vamathevan J., Bass S., Qin H., Dragoi I., Sellers P.,
RA McDonald L., Utterback T., Fleischmann R.D., Nierman W.C., White O.,

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RA Salzberg S.L., Smith H.O., Colwell R.R., Mekalanos J.J., Venter J.C.,
 RA Fraser C.M.;
 RT "DNA sequence of both chromosomes of the cholera pathogen *Vibrio*
 RT *cholerae*.";
 RL Nature 406:477-483(2000).
 CC -1- CATALYTIC ACTIVITY: GLUCOSE 6-PHOSPHATE = FRUCTOSE 6-PHOSPHATE.
 CC -1- PATHWAY: INVOLVED IN GLYCOLYSIS AND IN GLUCONEOGENESIS.
 CC -1- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
 CC -1- SIMILARITY: BELONGS TO THE GPI FAMILY.
 CC -----
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 CC -----
 DR EMBL; AE004125; AAF93547.1; -.
 DR TIGR; VC0374; -.
 DR InterPro; IPR001672; G6P_Isomerase.
 DR Pfam; PF00342; PCI_1.
 DR PRINTS; PR00562; G6PISOMERASE.
 DR PROSITE; PS00765; P-GLUCOSE-ISOMERASE_1; 1.
 DR PROSITE; PS00174; P-GLUCOSE-ISOMERASE_2; 1.
 KW Isomerase; Gluconeogenesis; Glycolysis; Complete proteome.
 FT ACT_SITE 387 BY SIMILARITY.
 FT ACT_SITE 515 BY SIMILARITY.
 SQ SEQUENCE 550 AA; 60690 MW; 5E3880421C3A1B16 CRC64;
 Query Match 69.1%; Score 38; DB 1; Length 550;
 Best Local Similarity 66.7%; Pred. NO. 4.7;
 Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;
 QY 1 HHQKLV--FFAE 10
 DB 417 HHQKLSNFFAQ 428
 RESULT 14
 SAS_PIG STANDARD; PRT; 109 AA.
 ID SAS_PIG
 AC Q29257;
 DT 15-JUL-1999 (Rel. 38, Created)
 DT 15-JUL-1999 (Rel. 38, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Sarcoma amplified sequence (Fragment).
 GN SAS.
 OS *Sus scrofa* (Pig).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
 OX NCBI_TaxID=9823;
 RN [1]
 RP SEQUENCE FROM N.A.
 RT TISSUE=Small intestine;
 RX MEDLINE=96327607; PubMed=8672129;
 RA Winteroe A.K., Fredholm M., Davies W.;
 RT "Evaluation and characterization of a porcine small intestine cDNA
 RT library: analysis of 839 clones.";
 RL Mamm. Genome 7:509-517(1996).
 CC -1- SUBCELLULAR LOCATION: Integral membrane protein.
 CC -1- SIMILARITY: BELONGS TO THE TETRASPANIN (TM4SF) FAMILY.
 DR InterPro; IPR000301; Transmem_4.
 DR Pfam; PF00335; transmembrane4; 1.
 DR PRINTS; PR00259; TMFOUR.
 KW Glycoprotein; Transmembrane.
 FT DOMAIN 1 12 CYTOPLASMIC (POTENTIAL).
 FT TRANSMEM 13 33 POTENTIAL.
 FT DOMAIN 34 44 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 45 65 POTENTIAL.
 FT DOMAIN 66 72 CYTOPLASMIC (POTENTIAL).
 FT TRANSMEM 73 93 POTENTIAL.
 FT DOMAIN 94 >109 EXTRACELLULAR (POTENTIAL).

FT CARBOHYD 100 100 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT NON_TER 109 109
 SQ SEQUENCE 109 AA; 11291 MW; 5CC5EAB8B7F152B1 CRC64;
 Query Match 65.5%; Score 36; DB 1; Length 109;
 Best Local Similarity 75.0%; Pred. NO. 2.2;
 Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;
 QY 1 HHQKLVFF 8
 DB 70 HHQVLEFF 77
 RESULT 15
 TNE6_HUMAN STANDARD; PRT; 204 AA.
 ID TNE6_HUMAN
 AC O95857;
 DT 16-OCT-2001 (Rel. 40, Created)
 DT 16-OCT-2001 (Rel. 40, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Tetraspan NET-6.
 GN TNE6.
 OS *Homo sapiens* (Human).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
 OX NCBI_TaxID=9606;
 RN [1]
 RP SEQUENCE FROM N.A.
 RT Rubinstein E., Serru V., Dessen P., Boucheix C.;
 RA "New tetraspans identified in the EST database.";
 RL Submitted (JAN-1999) to the EMBL/GenBank/DBJ databases.
 [2]
 RP SEQUENCE FROM N.A.
 RC TISSUE=Pituitary;
 RX MEDLINE=20402571; PubMed=10931946;
 RA Hu R.-W., Han Z.-G., Song H.-D., Peng Y.-D., Huang Q.-H., Ren S.-X.,
 RA Gu Y.-J., Huang C.-H., Li Y.-B., Jiang C.-L., Fu G., Zhang Q.-H.,
 RA Gu B.-W., Dai M., Mao Y.-F., Gao G.-F., Rong R., Ye M., Zhou J.,
 RA Xu S.-H., Gu J., Shi J.-X., Jin W.-R., Zhang C.-K., Wu T.-M.,
 RA Huang G.-Y., Chen Z., Chen M.-D., Chen J.-L.;
 RT "Gene expression profiling in the human hypothalamus-pituitary-adrenal
 RT axis and full-length cDNA cloning.";
 RL Proc. Natl. Acad. Sci. U.S.A. 97:9543-9548(2000).
 CC -1- SUBCELLULAR LOCATION: Integral membrane protein (Probable).
 CC -1- SIMILARITY: BELONGS TO THE TETRASPANIN (TM4SF) FAMILY.
 CC -----
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 CC -----
 DR EMBL; AF120265; AADI7294.1; -.
 DR EMBL; AF100759; AAD43023.1; -.
 DR InterPro; IPR000301; Transmem_4.
 DR Pfam; PF00335; transmembrane4; 1.
 DR PRINTS; PR00259; TMFOUR.
 DR PROSITE; PS00421; TM4_1; FALSE_NEG.
 KW Glycoprotein; Transmembrane.
 FT DOMAIN 1 19 CYTOPLASMIC (POTENTIAL).
 FT TRANSMEM 20 40 POTENTIAL.
 FT DOMAIN 41 44 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 45 65 POTENTIAL.
 FT DOMAIN 66 72 CYTOPLASMIC (POTENTIAL).
 FT TRANSMEM 73 93 POTENTIAL.
 FT DOMAIN 94 167 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 168 188 POTENTIAL.
 FT DOMAIN 189 204 CYTOPLASMIC (POTENTIAL).
 FT CARBOHYD 113 113 N-LINKED (GLCNAC. . .) (POTENTIAL).
 FT TRANSMEM 137 137 N-LINKED (GLCNAC. . .) (POTENTIAL).
 SQ SEQUENCE 204 AA; 22147 MW; 5928646BCD83C0D6 CRC64;


```
RESULT 18
G6PI_PASMU STANDARD; PRT; 549 AA.
ID Q9CNL2: 2002 (Rel. 41, Created)
AC Q9CNL2: 2002 (Rel. 41, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Glucose-6-phosphate isomerase (EC 5.3.1.9) (GPI) (Phosphoglucose
DE isomerase) (PGI) (Phosphohexose isomerase) (PHI).
GN PGI OR PM0416.
OS Pasteurella multocida.
OC Bacteria; Proteobacteria; gamma subdivision; Pasteurellaceae;
OC Pasteurella.
OX NCBI_TaxID=747;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=PM70;
RX MEDLINE=21143866; PubMed=11248100;
RA May B.J., Zhang Q., Li L.L., Paustian M.L., Whittam T.S., Kapur V.;
RT "Complete genomic sequence of Pasteurella multocida Fm70.";
RL Proc. Natl. Acad. Sci. U.S.A. 98:3460-3465(2001).
CC -|- CATALYTIC ACTIVITY: GLUCOSE 6-PHOSPHATE -> FRUCTOSE 6-PHOSPHATE.
CC -|- PATHWAY: INVOLVED IN GLYCOLYSIS AND IN GLUCONEOGENESIS.
CC -|- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
CC -|- SIMILARITY: BELONGS TO THE GPI FAMILY.
-----
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DR EMBL; AE006077; AAK02500.1; -
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00562; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
DR PROSITE; PS00174; P_GLUCOSE_ISOMERASE_2; 1.
KW Isomerase; Gluconeogenesis; Glycolysis; Complete proteome.
FT ACT_SITE 387 BY SIMILARITY.
FT ACT_SITE 515 515
SQ SEQUENCE 549 AA; 61437 MW; E5E4856927B93283 CRC64;

Query Match 63.6%; Score 35; DB 1; Length 549;
Best Local Similarity 58.3%; Pred. No. 18;
Matches 7; Conservative 3; Mismatches 0; Indels 2; Gaps 1;

OY 1 HHOKLV--FFAE 10
DB 417 HHEKLSNFFAQ 428
||||: |||:

RESULT 19
RSG2_HUMAN STANDARD; PRT; 849 AA.
ID RSG2_HUMAN STANDARD; PRT; 849 AA.
AC Q15283: Q15284; Q00895; Q99577; Q92594; Q90EQ2;
DT 16-OCT-2001 (Rel. 40, Created)
DE Ras GTPase-activating protein 2 (GAPIM)
DE 16-OCT-2001 (Rel. 40, Last sequence update)
DE 16-OCT-2001 (Rel. 40, Last annotation update)
DE Ras GTPase-activating protein 2 (GAPIM).
GN RAS2 OR RASGAP OR GAPIM.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=97074668; PubMed=8917095;
RA Kobayashi M., Masui T., Kusuda J., Kameoka Y., Hashimoto K.,
RA Iwashita S.;

"Human rasGTPase-activating protein (human counterpart of GAP1m):
sequence of the cDNA, primary structure of the protein, production and
chromosomal localization.";
Gene 175:173-177(1996).
[2]
RN RP SEQUENCE FROM N.A.
RX TISSUE=Brain;
RX MEDLINE=97001173; PubMed=8812506;
RA Li S., Satoh H., Watanabe T., Nakamura S., Hattori S.;
RT "cDNA cloning and chromosomal mapping of a novel human GAP (GAP1m), a
RT GTPase-activating protein of Ras.";
RL Genomics 35:625-627(1996).
[3]
RN RP SEQUENCE FROM N.A.
RX TISSUE=Blood;
RX MEDLINE=98044291; PubMed=9382842;
RA Lockyer P.J., Bottomley J.R., Reynolds J.S., McNulty T.J.,
RA Venkateswarlu K., Potter B.V.L., Dempsey C.E., Cullen P.J.;
RT "Distinct subcellular localisations of the putative inositol 1,3,4,5-
RT tetrakisphosphate receptors GAP1(IP4BP) and GAP1m result from the
RT GAP1(IP4BP) PH domain directing plasma membrane targeting.";
RL Curr. Biol. 7:1007-1010(1997).
[4]
RN RP SEQUENCE FROM N.A.
RX TISSUE=Blood;
RA Lockyer P.J.;
Submitted (DEC-1998) to the EMBL/GenBank/DBJ databases.
CC -|- FUNCTION: INHIBITORY REGULATOR OF THE RAS-CYCLIC AMP PATHWAY.
CC -|- BINDS INOSITOL TETRAKISPHOSPHATE (IP4).
CC -|- SUBCELLULAR LOCATION: PERINUCLEAR AND CYTOPLASMIC.
CC -|- SIMILARITY: CONTAINS 2 C2 DOMAINS.
CC -|- SIMILARITY: CONTAINS 1 PH DOMAIN.
CC -|- SIMILARITY: CONTAINS 1 BTK DOMAIN.
CC -|- SIMILARITY: CONTAINS 1 RAS-GAP DOMAIN.
-----
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DR EMBL; D78155; BAAL1230.1; -
DR EMBL; D78156; BAAL1231.1; -
DR EMBL; D82880; BAAL1621.1; -
DR EMBL; D82881; BAAL1622.1; -
DR EMBL; AF115573; AAD09821.1; -
DR HSSP; P21707; IRSY.
DR MIN; 601589; -
DR InterPro; IPR001562; BTK.
DR InterPro; IPR000008; C2.
DR InterPro; IPR001849; PH.
DR InterPro; IPR001936; RasGAP.
DR Pfam; PF00779; BTK; 1.
DR Pfam; PF00168; C2; 2.
DR Pfam; PF00169; PH; 1.
DR Pfam; PF00616; RasGAP; 1.
DR PRINTS; PR00402; TECBTKDOMAIN.
DR SMART; SM00107; BTK; 1.
DR SMART; SM00239; C2; 2.
DR SMART; SM00233; PH; 1.
DR SMART; SM00323; RasGAP; 1.
DR PROSITE; PS50003; PH_DOMAIN; 1. FALSE_NEG.
DR PROSITE; PS00499; C2_DOMAIN_1; 2.
DR PROSITE; PS50004; C2_DOMAIN_2; 2.
DR PROSITE; PS00509; RAS_GTPASE_ACTIV_1; FALSE_NEG.
DR PROSITE; PS50018; RAS_GTPASE_ACTIV_2; 1.
KW GTPase activation; Repeat.
FT DOMAIN 25 122 C2 DOMAIN 1.
FT DOMAIN 166 273 C2 DOMAIN 2.
FT DOMAIN 356 550 RAS-GAP.
FT DOMAIN 604 705 PH.
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FT DOMAIN 707 743 BTK.
FT DOMAIN 1 26 ALA-RICH.
FT CONFLICT 216 216 T -> A (IN REF. 1).
FT CONFLICT 645 645 G -> GS (IN REF. 1).
FT CONFLICT 645 645 G -> EFIER (IN REF. 2).
SQ SEQUENCE 849 AA: 96526 MW: A4B49IDFF5C4CB76 CRC64;

Query Match 63.6%; Score 35; DB 1; Length 849;
Best Local Similarity 77.8%; Pred. No. 29;
Matches 7; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHOKLVFFA 9
  || || || ||
DB 370 HHDKLVFFA 378

RESULT 20
KROS_HUMAN STANDARD; PRT: 2347 AA.
ID KROS_HUMAN Q15368;
AC P08922; Q15368;
DT 01-NOV-1988 (Rel. 09, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 01-NOV-1997 (Rel. 35, Last annotation update)
DE Proto-oncogene tyrosine-protein kinase ROS precursor (EC 2.7.1.112).
GN ROS1.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=90280463; PubMed=2352949;
RA Birchmeier C., O'Neill K., Riggs M., Wigler M.;
RT "Characterization of ROS1 cDNA from a human glioblastoma cell line.";
RL Proc. Natl. Acad. Sci. U.S.A. 87:4799-4803(1990).
RN [2]
RP SEQUENCE OF 1790-2259 FROM N.A.
RX MEDLINE=87064611; PubMed=3023956;
RA Matsushima H., Wang L.-H., Shibuya M.;
RT "Human c-ros-1 gene homologous to the v-ros sequence of UR2 sarcoma
  virus encodes for a transmembrane receptorlike molecule.";
RL Mol. Cell. Biol. 6:3000-3004(1986).
RN [3]
RP SEQUENCE OF 1854-2245 FROM N.A.
RX MEDLINE=87064625; PubMed=3785223;
RA Birchmeier C., Birnbaum D., Waitches G., Fasano O., Wigler M.;
RT "Characterization of an activated human ros gene.";
RL Mol. Cell. Biol. 6:3109-3116(1986).
CC -!- FUNCTION: THIS IS A PROBABLY A CELL GROWTH OR DIFFERENTIATION
  FACTOR RECEPTOR WITH A TYROSINE-PROTEIN KINASE ACTIVITY.
CC -!- CATALYTIC ACTIVITY: ATP + a protein tyrosine = ADP + protein
  tyrosine phosphate.
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: BELONGS TO THE INSULIN RECEPTOR FAMILY OF TYROSINE-
  PROTEIN KINASES. SEVENLESS SUBFAMILY.
CC -----
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  or send an email to license@isb-sib.ch).
CC -----
DR EMBL: M34353; AAA60278.1;
DR EMBL: M13599; AAA60277.1;
DR EMBL: M13368; AAA60277.1; JOINED.
DR EMBL: M13591; AAA60277.1; JOINED.
DR EMBL: M13592; AAA60277.1; JOINED.
DR EMBL: M13593; AAA60277.1; JOINED.
DR EMBL: M13594; AAA60277.1; JOINED.
DR EMBL: M13595; AAA60277.1; JOINED.
DR EMBL: M13596; AAA60277.1; JOINED.

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DR EMBL: M13597; AAA60277.1; JOINED.
DR EMBL: M13598; AAA60277.1; JOINED.
DR EMBL: M13880; AAA36580.1; ALT_TERM.
DR PIR: A25223; TVHURS.
DR PIR: A24421; TVHURT.
DR HSP: P08631; IAD5.
DR MIM: 165020;
DR InterPro: IPR000719; Euk_pkinase.
DR InterPro: IPR003961; FN.III.
DR InterPro: IPR000033; Ldl_receptor_rep.
DR InterPro: IPR002011; Receptor_tyr_kin_II.
DR InterPro: IPR001245; Tyr_pkinase.
DR Pfam: PF00041; fn3; 7.
DR Pfam: PF00069; pkinase; 1.
DR SMART: SM00060; FN3; 5.
DR SMART: SM00135; LY; 2.
DR SMART: SM00219; TyrcK; 1.
DR PROSITE: PS00107; PROTEIN_KINASE_ATP; 1.
DR PROSITE: PS00109; PROTEIN_KINASE_TYR; 1.
DR PROSITE: PS00239; RECEPTOR_TYR_KIN_II; 1.
DR PROSITE: PS00111; PROTEIN_KINASE_DOM; 1.
DR Transfaser: Tyrosine-protein kinase; Receptor; Transmembrane;
KW Glycoprotein; ATP-binding; Phosphorylation; Proto-oncogene;
KW Signal.
FT SIGNAL 1 27 POTENTIAL.
FT CHAIN 28 2347 PROTO-ONCOGENE TYROSINE-PROTEIN KINASE
  ROS.
FT DOMAIN 28 1859 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 1860 1882 POTENTIAL.
FT DOMAIN 1883 2347 CYTOPLASMIC (POTENTIAL).
FT DOMAIN 1945 2222 PROTEIN KINASE.
FT NP_BIND 1951 1959 ATP (BY SIMILARITY).
FT BINDING 1980 1980 ATP (BY SIMILARITY).
FT MOD_RES 2114 2114 PHOSPHORYLATION (AUTO-) (BY SIMILARITY).
FT CARBOHYD 52 52 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 114 114 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 123 123 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 324 324 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 352 352 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 396 396 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 471 471 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 607 607 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 628 628 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 706 706 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 714 714 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 732 732 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 939 939 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 961 961 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1015 1015 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1087 1087 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1090 1090 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1095 1095 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1211 1211 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1272 1272 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1330 1330 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1458 1458 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1461 1461 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1474 1474 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1499 1499 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1565 1565 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1669 1669 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1715 1715 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1738 1738 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 1808 1808 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CONFLICT 2213 2213 N -> D (IN REF. 2 AND 3).
FT CONFLICT 2228 2228 QC -> KS (IN REF. 2 AND 3).
FT CONFLICT 2246 2246 EDGDVCLNSDDIM -> KFSSEFSFRCVTN
  (IN REF. 2).
SQ SEQUENCE 2347 AA: 263956 MW: E14F3DFD410C1D2A CRC64;

```

Query Match 63.6%; Score 35; DB 1; Length 2347;
 Best Local Similarity 55.6%; Pred. No. 81;
 Matches 5; Conservative 4; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQKLVFAE 10
|:|:|:|:
Db 333 HQQIVFESE 341

RESULT 21

SLP_BACSU STANDARD; PRT; 124 AA.
AC P39910;
DT 01-FEB-1995 (Rel. 31, Created)
DT 01-FEB-1995 (Rel. 31, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE PAL-related lipoprotein precursor.
GN SLP OR PAL.
OS Bacillus subtilis.
OC Bacteria; Firmicutes; Bacillus/Clostridium group;
OC Bacillus/Staphylococcus group; Bacillus.
OX NCBI_TaxID=1423;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=168;
RX MEDLINE=90368558; PubMed=1697575;
RA Hemilae H.O., Palva A., Paulin L., Arvidson S., Palva I.;
RT "Secretory S complex of Bacillus subtilis: sequence analysis and
RT identity to pyruvate dehydrogenase."; J. Bacteriol. 172:5052-5063(1990).
RL [2]
RN SEQUENCE FROM N.A.
RP STRAIN=168;
RC STRAIN=168;
RX MEDLINE=92038903; PubMed=1936936;
RA Hemilae H.;
RT "Sequence of a PAL-related lipoprotein from Bacillus subtilis."; FEMS Microbiol. Lett. 66:37-41(1991).
RN [3]
RP SEQUENCE FROM N.A.
RC STRAIN=168;
RX MEDLINE=97124187; PubMed=8969500;
RA Winters P., Caldwell R.M., Enfield L., Ferrari E.;
RT "The ampS-nprE (124 degrees-127 degrees) region of the Bacillus
RT subtilis 168 chromosome: sequencing of a 27 kb segment and
RT identification of several genes in the area."; Microbiology 142:3033-3037(1996).
RN [4]
RP SEQUENCE FROM N.A.
RC STRAIN=168;
RA Caldwell R.M., Ferrari E.;
RT "Sequence analysis of the mobA-ampS region of the Bacillus subtilis
RT chromosome."; Submitted (JUL-1997) to the EMBL/GenBank/DBJ databases.
CC -|- SUBCELLULAR LOCATION: Attached to the membrane by a lipid anchor
CC (Probable).
CC
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CC
CC EMBL; M57435; AAA62685.1; -;
CC EMBL; AF012285; AAC24936.1; -;
CC EMBL; Z99111; CAB13335.1; -;
CC PIR; B54546; B54546.
CC Subtilisin; BG10211; slp.
DR PROSITE; PS00013; PROKAR_LIPOPROTEIN; 1.
KW Membrane; Lipoprotein; Signal; Complete proteome.
FT SIGNAL 1 18 POTENTIAL.
FT CHAIN 19 124 PAL-RELATED LIPOPROTEIN.
FT LIPID 19 19 N-ACYL DIGLYCERIDE (PROBABLE).
SQ SEQUENCE 124 AA; 14538 MW; 804401AF0E88446F CRC64;

Query Match 61.8%; Score 34; DB 1; Length 124;
Best Local Similarity 40.0%; Pred. No. 6.3;
Matches 4; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFAE 10
|:|:|:|:
Db 36 HHTQILFFSD 45

RESULT 22

YNSI_YEAST STANDARD; PRT; 763 AA.
AC P42843;
DT 01-NOV-1995 (Rel. 32, Created)
DT 01-NOV-1995 (Rel. 32, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Hypothetical 89.9 kDa protein in RPA2-StBI intergenic region.
GN YNL311C OR N0376.
OS Saccharomyces cerevisiae (Baker's yeast).
OC Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
OC Saccharomycetales; Saccharomycetaceae; Saccharomyces.
OX NCBI_TaxID=4932;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=S288C / FY1676;
RX MEDLINE=96076632; PubMed=7502583;
RA Maftahi M., Nicaud J.-M., Levesque H., Gaillardin C.;
RT "Sequencing analysis of a 24.7 kb fragment of yeast chromosome XIV
RT identifies six known genes, a new member of the hexose transporter
RT family and ten new open reading frames."; Yeast 11:1077-1085(1995).
RN [2]
RP SEQUENCE OF 149-763 FROM N.A.
RA Maurer C.T.C., Urbanus J.H.M., Planta R.J.;
RL Submitted (MAY-1996) to the EMBL/GenBank/DBJ databases.
CC -|- SIMILARITY: CONTAINS 1 F-BOX DOMAIN.
CC
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CC
CC EMBL; Z46259; CA86384.1; -;
CC EMBL; Z71587; CA96240.1; -;
CC SGD; S0005255; YNL311C.
DR InterPro; IPR001810; F-box.
DR Pfam; PF00646; F-box; 1.
DR SMART; SM00256; FBOX; 1.
DR PROSITE; PS0181; FBOX; 1.
KW Hypothetical protein.
FT DOMAIN 54 100 F-BOX.
FT DOMAIN 22 28 POLY-GLU.
SQ SEQUENCE 763 AA; 88941 MW; 81102168449051BC CRC64;
Query Match 61.8%; Score 34; DB 1; Length 763;
Best Local Similarity 100.0%; Pred. No. 41;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
|:|:|:|:
Db 323 HHQKLV 328

RESULT 23
RPOD_PEA STANDARD; PRT; 1163 AA.
AC P12227;
DT 01-OCT-1989 (Rel. 12, Created)
DT 01-OCT-1989 (Rel. 12, Last sequence update)
DT 01-JUL-1993 (Rel. 26, Last annotation update)

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DE DNA-directed RNA polymerase beta" chain (EC 2.7.7.6) (Fragment).
GN RPOC2.
OS Pisum sativum (Garden pea).
OG Chloroplast.
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
OC eurosids I; Fabales; Fabaceae; Papilionoideae; Viciaeae; Pisum.
OX NCBI_TaxID=3888;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=86323089; PubMed=3530249;
RA Cozens A.L., Walker J.E.;
RT "Pea chloroplast DNA encodes homologues of Escherichia coli ribosomal
RT subunit S2 and the beta'-subunit of RNA polymerase.";
RL Biochem. J. 236:453-460(1986).
CC -!- FUNCTION: DNA-DEPENDENT RNA POLYMERASE CATALYZES THE TRANSCRIPTION
CC OF DNA INTO RNA USING THE FOUR RIBONUCLEOSIDE TRIPHOSPHATES AS
CC SUBSTRATES.
CC -!- CATALYTIC ACTIVITY: N nucleoside triphosphate = N diphosphate +
CC {RNA}(N).
CC -!- SUBUNIT: IN CHLOROPLAST THE RNA POLYMERASE IS COMPOSED OF FOUR
CC SUBUNITS: ALPHA, BETA, BETA', AND BETA".
CC -----
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CC -----
DR EMBL; X03912; CAA27545.1; -.
DR PIR; S07137; S07137.
DR Mendel; 5368; PISsa:rpoc2.1.
KW Transferase; Transcription; DNA-directed RNA polymerase; Chloroplast.
FT NON_TER 1
SQ SEQUENCE 1163 AA; 133598 MW; C92E7BE0A3FDB525 CRC64;

Query Match 61.8%; Score 34; DB 1; Length 1163;
Best Local Similarity 66.7%; Pred. No. 62;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Oy 1 HHQKLVFFA 9
Db 1149 HHRKLLDFA 1157

RESULT 24
ROM2_YEAST
ID ROM2_YEAST STANDARD; PRT; 1356 AA.
AC P51862;
DT 01-OCT-1996 (Rel. 34, Created)
DT 01-OCT-1996 (Rel. 34, Last sequence update)
DE 01-MAR-2002 (Rel. 41, Last annotation update)
DE RH01 GDP-GTP exchange protein 2.
GN ROM2 OR YLR371W OR L8039.3.
OS Saccharomyces cerevisiae (Baker's yeast).
OC Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
OC Saccharomycetales; Saccharomycetaceae; Saccharomycetes.
OX NCBI_TaxID=4932;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=S288C / AB972;
RA Johnston M., Andrews S., Brinkman R., Cooper J., Ding H., Du Z.,
RA Favell A., Fulton L., Gattung S., Greco T., Kirsten J.,
RA Kucaba T., Hallsworth K., Hawkins J., Hillier L., Jier M.,
RA Johnson D., Johnston L., Langston Y., Latreille P., Le T.,
RA Mardis E., Menezes S., Miller N., Nhan M., Pauley A., Peluso D.,
RA Rifken L., Riles L., Raich A., Trevaskis E., Vignati D.,
RA Wilcox L., Wohlman P., Vaudin M., Wilson R., Waterston R.;
RL Submitted (JAN-1995) to the EMBL/GenBank/DBJ databases.
RN [2]
RP CHARACTERIZATION.

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RX MEDLINE=96208506; PubMed=8641285;
RA Ozaki K., Tanaka K., Imamura H., Hihara T., Kameyama T.,
RA Nonaka H., Hirano H., Matsuura Y., Takai Y.;
RT "Romlp and Rom2p are GDP/GTP exchange proteins (GEPs) for the Rhop1
RT small GTP binding protein in Saccharomyces cerevisiae.";
RL EMBO J. 15:2196-2207(1996).
CC -!- FUNCTION: STIMULATES THE EXCHANGE OF RH01 GDP-BOUND FORM INTO
CC GTP-BOUND FORM.
CC -!- SIMILARITY: CONTAINS 1 DBL-HOMOLOGY DOMAIN (DH).
CC -----
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CC -----
DR EMBL; U19103; AAB67564.1; -.
DR SGD; S0004363; ROM2.
DR InterPro; IPR001180; CNH.
DR InterPro; IPR000591; DEP.
DR InterPro; IPR000219; RhoGEF.
DR Pfam; PF00780; CNH; 1.
DR Pfam; PF00610; DEP; 1.
DR Pfam; PF00621; RhoGEF; 1.
DR SMART; SM00036; CNH; 1.
DR SMART; SM00049; DEP; 1.
DR SMART; SM00325; RhoGEF; 1.
DR PROSITE; PS00741; DH_1; FALSE_NEG.
DR PROSITE; PS50010; DH_2; 1.
KW Guanine-nucleotide releasing factor.
FT DOMAIN 659 846 DH.
FT DOMAIN 252 265 POLY-ASN.
FT DOMAIN 329 336 POLY-HIS.
FT DOMAIN 632 635 POLY-ASP.
SQ SEQUENCE 1356 AA; 152595 MW; 5FBC542114E7BC92 CRC64;

Query Match 61.8%; Score 34; DB 1; Length 1356;
Best Local Similarity 50.0%; Pred. No. 73;
Matches 7; Conservative 3; Mismatches 0; Indels 4; Gaps 1;

Oy 1 HHQKLV----FFAE 10
Db 1131 HKKELINHVFFAE 1144

RESULT 25
TRX2_HUMAN
ID TRX2_HUMAN STANDARD; PRT; 2715 AA.
AC Q9UMN6; Q9UK25; Q95836; Q9Y669; Q9Y668; O15022;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Trithorax homolog 2 (Mixed lineage leukemia gene homolog 2 protein).
GN TRX2 OR HRX2 OR MLL2 OR KIAA0340.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A. (LONG ISOFORM).
RA Angrand P.O., Valvatne H., Jeanmougin F., Adamson A.,
RA van der Hoeven F., Olsen L., Tekotte H., Huang N., Poch O.,
RA Lamerdin J., Chambon P., Lossen R., Stewart A., Aasland R.;
RT "Mammalian trithorax- and ASH1-like proteins: putative chromatin
RT regulators which contain PHD fingers and SET domains.";
RL Submitted (JUN-1998) to the EMBL/GenBank/DBJ databases.
RN [2]
RP SEQUENCE FROM N.A. (LONG ISOFORM).
RA Lamerdin J.E., McCready P.M., Adamson A.W., Burkhardt-Schultz K.,
RA Garcia E., Kyle A., Ramirez M., Stillwagen S., Garnes J., Danganan L.,
RA Bruce R., Quan G., Montgomery M., Ow D., Kobayashi A., Olsen A.O.,

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CC THE SYNTHESIS OF A POLYKETIDE MOLECULE WHICH MAY BE INVOLVED IN
CC SECONDARY METABOLISM.
CC -!- COFACTOR: CONTAINS 5 COVALENTLY BOUND PHOSPHOPANTETHEINES
CC (POTENTIAL).
CC -!- SIMILARITY: CONTAINS 5 ACYL CARRIER DOMAINS.
CC -----
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CC -----
DR EMBL; Z14098; CAA78479.1; -
DR EMBL; U11039; AAB85145.1; -
DR EMBL; Z35133; CAA84504.1; -
DR EMBL; Z99113; CAB13602.1; -
DR PIR; S25021; S25021.
DR HSSP; P27796; LPXT.
DR Subtilist; BG10698; pksL.
DR InterPro; IPR000794; Ketoacyl-synt.
DR InterPro; IPR003880; Phosphopant_attach.
DR Pfam; PF00109; ketoacyl-synt; 4.
DR Pfam; PF02801; ketoacyl-synt_C; 4.
DR Pfam; PF00550; pp-binding; 5.
DR PROSITE; PS00012; PHOSPHOPANTETHEINE; 5.
DR PROSITE; PS00606; B.KETOACYL_SYNTHASE; 1.
DR PROSITE; PS00075; ACP_DOMAIN; 5.
DR Transferase; Acyltransferase; Antibiotic biosynthesis; NADP;
KW Phosphopantetheine; Multifunctional enzyme; Repeat; Complete proteome.
KW Phosphopantetheine; Multifunctional enzyme; Repeat; Complete proteome.
FT DOMAIN 211 280
FT DOMAIN 382 759
FT DOMAIN 937 1115
FT DOMAIN 1409 1602
FT DOMAIN 1687 1759
FT DOMAIN 1876 2253
FT DOMAIN 2491 2560
FT DOMAIN 2632 2701
FT DOMAIN 2823 3182
FT DOMAIN 3575 3776
FT DOMAIN 3854 3923
FT DOMAIN 4019 4373
FT BINDING 243 243
FT BINDING 1723 1723
FT BINDING 2523 2523
FT BINDING 2664 2664
FT BINDING 3886 3886
FT BINDING 3886 3886
SQ SEQUENCE 4427 AA; 493398 MW; 9612521E561AB9F2 CRC64;

Query Match 61.8%; Score 34; DB 1; Length 4427;
Best Local Similarity 100.0%; Pred. No. 2.4e+02;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
DB 691 HHQKLV 696
|||||

RESULT 27
ACUB_BACSU STANDARD; PRT; 214 AA.
AC P39066;
DT 01-FEB-1995 (Rel. 31, Created)
DT 01-FEB-1995 (Rel. 31, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Acetoin utilization acbB protein.
GN ACUB.
OS Bacillus subtilis.
OC Bacteria; Firmicutes; Bacillus/Clostridium group;
OC Bacillus/Staphylococcus group; Bacillus.
OX NCBI_TaxID=1423;
RN [1]
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RP SEQUENCE FROM N.A.
RC STRAIN=168;
RX MEDLINE=95020526; PubMed=7934817;
RA Grundy F.J., Waters D.A., Takova T.Y., Henkin T.M.;
RT "Identification of genes involved in utilization of acetate and
RL Mol. Microbiol. 10:259-271(1993).
RN [2]
RP SEQUENCE FROM N.A.
RX MEDLINE=98048467; PubMed=9387221;
RA Lapidus A., Galleron N., Sorokin A., Ehrlich S.D.;
RT "Sequencing and functional annotation of the Bacillus subtilis genes
RL in the 200 kb rrbB-dnaB region.";
RL Microbiology 143:3431-3441(1997).
CC -!- FUNCTION: ROLE IN GROWTH AND SPOULATION ON ACETOIN OR BUTANEDIOL.
CC INVOLVED IN THE BREAKDOWN OF THESE COMPOUNDS USED AS A CARBON
CC SOURCE.
CC -!- SIMILARITY: CONTAINS 2 CBS DOMAINS.
CC -----
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CC -----
DR EMBL; L17309; AAA68285.1; -
DR EMBL; AF008220; AAC00395.1; -
DR EMBL; Z99119; CAB14948.1; -
DR PIR; S39644; S39644.
DR Subtilist; BG10368; acbB.
DR InterPro; IPR002912; ACT.
DR InterPro; IPR000644; CBS.
DR Pfam; PF01842; ACT; 1.
DR Pfam; PF00571; CBS; 2.
DR SMART; SM00116; CBS; 2.
KW Sporulation; Repeat; CBS domain; Complete proteome.
FT DOMAIN 5 58
FT DOMAIN 75 128
FT SEQUENCE 214 AA; 24351 MW; 3B7A964B5C95CCEFCRC64;

Query Match 60.0%; Score 33; DB 1; Length 214;
Best Local Similarity 83.3%; Pred. No. 17;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
DB 110 HHQKLV 115
|||||

RESULT 28
PPNK_THEME STANDARD; PRT; 258 AA.
ID PPNK_THEME
AC Q9X255;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Probable inorganic polyphosphate/ATP-NAD kinase (EC 2.7.1.23)
DE (poly(P)/ATP NAD kinase).
GN PPNK OR TM1733.
OS Thermotoga maritima.
OC Bacteria; Thermotogales; Thermotoga.
OX NCBI_TaxID=2336;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=MSB8 / DSM 3109;
RX MEDLINE=99287316; PubMed=10360571;
RA Nelson K.E., Clayton R.A., Gill S.R., Gwinn M.L., Dodson R.J.,
RA Haft D.H., Hickey E.K., Peterson J.D., Nelson W.C., Ketchum K.A.,
RA McDonald L., Utterback T.R., Malek J.A., Linher K.D., Garrett M.M.,
RA Stewart A.M., Cotton M.D., Pratt M.S., Phillips C.A., Richardson D.,
RA Heidelberg J., Sutton G.G., Fleischmann R.D., Eisen J.A., White O.,
```

RA Salzberg S.L., Smith H.O., Venter J.C., Fraser C.M.;
 RT "Evidence for lateral gene transfer between Archaea and Bacteria from
 RL genome sequence of *Thermotoga maritima*.";
 CC Nature 399:323-329(1999).
 CC -!- FUNCTION: Catalyzes the phosphorylation of NAD to NADP. Utilizes
 CC ATP and other nucleoside triphosphates as well as inorganic
 CC polyphosphate as a source of phosphorus (By similarity).
 CC -!- CATALYTIC ACTIVITY: ATP + NAD(+) = ADP + NADP(+).
 CC -!- COFACTOR: Requires divalent metal ions for activity (By
 CC similarity).
 CC -!- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
 CC -!- SIMILARITY: BELONGS TO THE NAD KINASE FAMILY.
 CC
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 CC
 CC EMBL: AF001812; AAC36798.1; -
 CC TIGR: TM1733; -
 CC InterPro: IPR002504; DUF15.
 CC Pfam: PF01513; DUF15; 1.
 CC Transferase: Kinase; NADP; Complete proteome.
 CC SEQUENCE 258 AA; 29241 MW; 45EBBCA0B6FD3EAB CRC64;
 CC
 CC Query Match 60.0%; Score 33; DB 1; Length 258;
 CC Best Local Similarity 50.0%; Pred. No. 21;
 CC Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;
 CC
 CC QY 1 HHOKLVFFAE 10
 CC || : ||:
 CC Db 140 HHSSWFFPAD 149
 CC
 CC RESULT 29
 CC IL0S_HUMAN
 CC ID IL0S_HUMAN STANDARD; PRT; 325 AA.
 CC AC Q08334;
 CC DT 01-FEB-1995 (Rel. 31, Created)
 CC DT 01-FEB-1995 (Rel. 31, Last sequence update)
 CC DT 01-MAR-2002 (Rel. 41, Last annotation update)
 CC DE Interleukin-10 receptor beta chain precursor (IL-10R-B) (IL-10R2)
 CC DE (Cytokine receptor class-II CR2-4).
 CC GN IL10RB OR CRFB4.
 CC OS Homo sapiens (Human).
 CC OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 CC OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
 CC OX NCBI_TaxID=9606;
 CC RN [1]
 CC RP SEQUENCE FROM N.A.
 CC RC TISSUE=Fetal brain;
 CC RX MEDLINE=93300510; PubMed=8314576;
 CC RA Lutfalla G., Gardiner K., Uze G.;
 CC RT "A new member of the cytokine receptor gene family maps on chromosome
 CC RT 21 at less than 35 kb from IFNAR.";
 CC RL Genomics 16:366-373(1993).
 CC RN [2]
 CC RP SEQUENCE FROM N.A.
 CC RX MEDLINE=96054036; PubMed=7563119;
 CC RA Lutfalla G., McInnis M.G., Antonarakis S.E., Uze G.;
 CC RT "Structure of the human CRFB4 gene: comparison with its IFNAR
 CC RT neighbor.";
 CC RL J. Mol. Evol. 41:338-344(1995).
 CC RN [3]
 CC RP CHARACTERIZATION.
 CC RX MEDLINE=97459974; PubMed=9312047;
 CC RA Kotenko S.V., Krause C.D., Iztocova L.S., Pollack B.P., Wu W.,
 CC RA Pestka S.;
 CC RT "Identification and functional characterization of a second chain of
 CC RT the interleukin-10 receptor complex.";

RL EMBO J. 16:5894-5903(1997).
 RN [4]
 RN CHARACTERIZATION.
 RX MEDLINE=20469498; PubMed=10875937;
 RA Xie M.-H., Aggarwal S., Ho W.-H., Foster J., Zhang Z., Stinson J.,
 RA Wood W.I., Goddard A.D., Gurney A.L.;
 RT "Interleukin (IL)-22, a novel human cytokine that signals through the
 RT interferon receptor-related proteins CRF2-4 and IL-22R.";
 RL J. Biol. Chem. 275:31335-31339(2000).
 CC -!- FUNCTION: RECEPTOR FOR IL-10 AND IL-22. SERVES AS AN ACCESSORY
 CC CHAIN ESSENTIAL FOR THE ACTIVE IL-10 RECEPTOR COMPLEX AND TO
 CC INITIATE IL-10-INDUCED SIGNAL TRANSDUCTION EVENTS.
 CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
 CC -!- SIMILARITY: CONTAINS 2 FIBRONECTIN TYPE III-LIKE DOMAINS.
 CC -!- SIMILARITY: BELONGS TO THE CLASS II CYTOKINE FAMILY OF RECEPTORS.
 CC
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 CC
 CC EMBL: Z17227; CAA78933.1; -
 CC EMBL: U08988; AAA86872.1; -
 CC PIR: A47003; A47003.
 CC HSP: P13726; ITPH.
 CC MIM: 123889; -
 CC InterPro: IPR000282; Cytok_receptor_2.
 CC InterPro: IPR01187; Tissue_fac.
 CC Pfam: PF01108; Tissue_fac; 1.
 CC Receptor; Transmembrane; Glycoprotein; Signal.
 FT SIGNAL 1 19 POTENTIAL
 FT CHAIN 20 325 INTERLEUKIN-10 RECEPTOR BETA CHAIN.
 FT DOMAIN 20 220 EXTRACELLULAR (POTENTIAL).
 FT TRANSMEM 221 242 POTENTIAL.
 FT DOMAIN 243 325 CYTOPLASMIC (POTENTIAL).
 FT DOMAIN 113 205 FIBRONECTIN TYPE-III.
 FT DISULFID 65 74 BY SIMILARITY.
 FT DISULFID 185 209 BY SIMILARITY.
 FT CARBOHYD 49 49 N-LINKED (GLCNAC. .) (POTENTIAL).
 FT CARBOHYD 63 68 N-LINKED (GLCNAC. .) (POTENTIAL).
 FT CARBOHYD 102 102 N-LINKED (GLCNAC. .) (POTENTIAL).
 FT CARBOHYD 161 161 N-LINKED (GLCNAC. .) (POTENTIAL).
 FT CONFLICT 124 124 A -> D (IN REF. 2).
 FT CONFLICT 269 274 FLGHP -> VGRME (IN REF. 2).
 FT CONFLICT 274 325 MISSING (IN REF. 2).
 SQ SEQUENCE 325 AA; 37011 MW; 66706C79F8514B23 CRC64;
 CC
 CC Query Match 60.0%; Score 33; DB 1; Length 325;
 CC Best Local Similarity 55.6%; Pred. No. 27;
 CC Matches 5; Conservative 2; Mismatches 2; Indels 0; Gaps 0;
 CC
 CC QY 1 HHOKLVFFA 9
 CC || : ||:
 CC Db 274 HHNTLLFFS 282
 CC
 CC RESULT 30
 CC C7D9_SOYBN
 CC ID C7D9_SOYBN STANDARD; PRT; 496 AA.
 CC AC O81971;
 CC DT 15-DEC-1998 (Rel. 37, Created)
 CC DT 15-DEC-1998 (Rel. 37, Last sequence update)
 CC DT 15-DEC-1998 (Rel. 37, Last annotation update)
 CC DE Cytochrome P450 71D9 (EC 1.14.-.-) (P450 CP3).
 CC GN CYP71D9
 CC OS Glycine max (Soybean).
 CC OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
 CC OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
 CC OC eurosids I; Fabales; Fabaceae; Papilionoideae; Phaseoleae; Glycine.
 CC OX NCBI_TaxID=3847;


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RN  SEQUENCE FROM N.A.
RP  STRAIN=CV. HARSOY 63;
RC  MEDLINE=98311068; PubMed=9648734;
RA  Schopfer C.R., Ebel J.;
RT  "Identification of elicitor-induced cytochrome P450s of soybean
RL  (Glycine max L.) using differential display of mRNA.";
RM  Mol. Gen. Genet. 258:315-322(1998).
CC  -!- SIMILARITY: BELONGS TO THE CYTOCHROME P450 FAMILY.
CC  -----
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CC  modified and this statement is not removed. Usage by and for commercial
CC  entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC  or send an email to license@isb-sib.ch).
CC  -----
DR  EMBL; Y10490; CAA71514.1; -;
DR  InterPro; IPR001128; Cyt_P450.
DR  Pfam; PF00067; P450; 1.
DR  PROSITE; PS00086; CYTOCHROME_P450; 1.
KW  Oxidoreductase; Monooxygenase; Heme.
FT  BINDING 437 437 HEME (BY SIMILARITY).
SQ  SEQUENCE 496 AA; 56205 MW; 9C90947C8A546CE1 CRC64;

Query Match 60.0%; Score 33; DB 1; Length 496;
Best Local Similarity 83.3%; Pred. No. 41;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
Db 197 HHQKLI 202
|||||
DE 16-OCT-2001 (Rel. 41, Last annotation update)
DE Threonyl-tRNA synthetase (EC 6.1.1.3) (Threonine--tRNA ligase)
DE (Thrs).
DE THRS OR MG375.
OS Mycoplasma genitalium.
OC Bacteria; Firmicutes; Bacillus/Clostridium group; Mollicutes;
OC Mycoplasmataceae; Mycoplasma.
OX NCBI_TaxID=2097;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=RD / KW20 / ATCC 51907;
RX MEDLINE=95350630; PubMed=7542800;
RA Fleischmann R.D., Adams M.D., White O., Clayton R.A., Kirkness E.F.,
RA Kervatage A.R., Bult C.J., Tomb J.-F., Dougherty B.A., Merrick J.M.,
RA McKenney K., Sutton G., Fitzhugh W., Fields C.A., Gocayne J.D.,
RA Scott J.D., Shirley R., Liu L.-I., Glodek A., Kelley J.M.,
RA Weidman J.F., Phillips C.A., Spriggs T., Hedblom E., Cotton M.D.,
RA Utterback T.R., Hanna M.C., Nguyen D.T., Saudek D.M., Brandon R.C.,
RA Fine L.D., Fritchman J.L., Fuhrmann J.L., Geoghagen N.S.M.,
RA Gnehm C.L., McDonald L.A., Small K.V., Fraser C.M., Smith H.O.,
RA Venter J.C.;
RT "Whole-genome random sequencing and assembly of Haemophilus
RT influenzae Rd.";
RL Science 269:496-512(1995).
CC -!- CATALYTIC ACTIVITY: D-glucose 6-phosphate = D-fructose 6-
CC phosphate.
CC -!- PATHWAY: INVOLVED IN GLYCOLYSIS AND IN GLUCONEOGENESIS.
CC -!- SUBCELLULAR LOCATION: Cytoplasmic (BY SIMILARITY).
CC -!- SIMILARITY: BELONGS TO THE GPI FAMILY.
CC -----
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CC or send an email to license@isb-sib.ch).
CC -----
DR  EMBL; U32831; AAC23219.1; ALT_INIT.
DR  TIGR; H11576; -;
DR  InterPro; IPR001672; G6P_Isomerase.
DR  Pfam; PF00342; PGI; 1.
DR  PRINTS; PR00662; G6PISOMERASE.
DR  PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
DR  PROSITE; PS00174; P_GLUCOSE_ISOMERASE_2; 1.
KW  Isomerase; Gluconeogenesis; Glycolysis; Complete proteome.
FT  ACT_SITE 387 387 BY SIMILARITY.
FT  ACT_SITE 515 515 BY SIMILARITY.
SQ  SEQUENCE 549 AA; 61622 MW; F65348C667068F16 CRC64;

Query Match 60.0%; Score 33; DB 1; Length 549;
Best Local Similarity 58.3%; Pred. No. 46;
Matches 7; Conservative 2; Mismatches 1; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
Db 417 HHNLLSNFFAQ 428
|||||
DE 01-FEB-1996 (Rel. 33, Created)
DE 01-FEB-1996 (Rel. 33, Last sequence update)
DE 16-OCT-2001 (Rel. 40, Last annotation update)
DE Threonyl-tRNA synthetase (EC 6.1.1.3) (Threonine--tRNA ligase)
DE (Thrs).
DE THRS OR MG375.
OS Mycoplasma genitalium.
OC Bacteria; Firmicutes; Bacillus/Clostridium group; Mollicutes;
OC Mycoplasmataceae; Mycoplasma.
OX NCBI_TaxID=2097;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=ATCC 33530 / G-37;
RX MEDLINE=96026346; PubMed=7569993;
RA Fraser C.M., Gocayne J.D., White O., Adams M.D., Clayton R.A.,
RA Fleischmann R.D., Bult C.J., Kervatage A.R., Sutton G., Kelley J.M.,
RA Fritchman J.L., Weidman J.F., Small K.V., Sandusky M., Fuhrmann J.L.,
RA Nguyen D.T., Utterback T.R., Saudek D.M., Phillips C.A., Merrick J.M.,
RA Tomb J.-F., Dougherty B.A., Bult K.F., Hu P.-C., Lucier T.S.,
RA Peterson S.N., Smith H.O., Hutchison C.A. III, Venter J.C.;
RT "The minimal gene complement of Mycoplasma genitalium.";
RL Science 270:397-403(1995).
RN [2]
RP SEQUENCE OF 350-463 FROM N.A.
RC STRAIN=ATCC 33530 / G-37;
RX MEDLINE=94075230; PubMed=8253680;
RA Peterson S.N., Hu P.-C., Bult K.F., Hutchison C.A. III;
RT "A survey of the Mycoplasma genitalium genome by using random
RT sequencing.";
RL J. Bacteriol. 175:7918-7930(1993).
CC -!- CATALYTIC ACTIVITY: ATP + L-threonine + tRNA(Thr) = AMP +
CC diphosphate + L-threonyl-tRNA(Thr).
CC -!- COFACTOR: BINDS 1 ZINC ION (BY SIMILARITY).
CC -!- SUBUNIT: HOMODIMER (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: Cytoplasmic.
CC -!- SIMILARITY: BELONGS TO CLASS-II AMINOACYL-TRNA SYNTHETASE FAMILY.
CC -----
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CC -----

DR EMBL; U39719; AAC71602.1; -

DR EMBL; U02130; AAD12408.1; -

DR HSSP; P00955; 1EVL.

DR TIGR; MG375; -

DR InterPro; IPR002106; AA_trna_ligase_II.

DR InterPro; IPR004154; HGTP_anticonodon.

DR InterPro; IPR002314; trna-synt_2b.

DR InterPro; IPR002320; trna-synt_thr.

DR Pfam; PF03129; HGTP_anticonodon; 1.

DR Pfam; PF00587; trna-synt_2b; 1.

DR PRINTS; PR01047; TRNASYNTHTHR.

DR PROSITE; PS00179; AA_TRNA_LIGASE_II_1; FALSE_NEG.

DR PROSITE; PS00339; AA_TRNA_LIGASE_II_2; FALSE_NEG.

KW Aminoacyl-trna synthetase; Protein biosynthesis; Ligase; ATP-binding;

KW Metal-binding; Zinc; Complete proteome.

FT DOMAIN 167 464 CATALYTIC.

FT METAL 260 260 ZINC (CATALYTIC) (BY SIMILARITY).

FT METAL 311 311 ZINC (CATALYTIC) (BY SIMILARITY).

FT METAL 441 441 ZINC (CATALYTIC) (BY SIMILARITY).

SQ SEQUENCE 564 AA; 65595 MW; 2CA833DA7F7AC447 CRC64;

Query Match 60.0%; Score 33; DB 1; Length 564;

Best Local Similarity 71.4%; Pred. No. 47;

Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHOKLVF 7

DB 207 HHQQLLF 213

RESULT 33

SDC3_CAEEL STANDARD; PRT; 2150 AA.

AC P34706;

DT 01-FEB-1994 (Rel. 28, Created)

DT 01-FEB-1994 (Rel. 28, Last sequence update)

DT 01-MAR-2002 (Rel. 41, Last annotation update)

DE Zinc finger protein sdc-3.

GN SDC-3.

OS Caenorhabditis elegans.

OS Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida; Rhabditoidea;

OC Rhabditidae; Peloderinae; Caenorhabditis.

OX NCBI_TaxID=6239;

RN [1]

RP SEQUENCE FROM N.A.

RC STRAIN-BRISTOL N2.

RX MEDLINE-93161411; PubMed=8431944;

RA Klein R.D., Meyer B.J.;

RT "Independent domains of the Sdc-3 protein control sex determination and dosage compensation in C. elegans."

RL Cell 72:349-364(1993).

CC -!- FUNCTION: CONTROLS BOTH SEX DETERMINATION AND X CHROMOSOME DOSAGE COMPENSATION. THESE TWO FUNCTIONS ACT INDEPENDENTLY.

CC -!- SUBCELLULAR LOCATION: Nuclear.

CC -!- DEVELOPMENTAL STAGE: EXPRESSED IN EMBRYONIC AND EARLY LARVAL STAGES.

CC -----

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CC -----

DR EMBL; M851149; AAA28144.1; -

DR PIR; S27802; S27802.

DR InterPro; IPR000822; Znf-C2H2.

DR PROSITE; PS00028; ZINC_FINGER_C2H2_1; 1.

DR PROSITE; PS50157; ZINC_FINGER_C2H2_2; FALSE_NEG.

KW Developmental protein; Zinc-finger; Metal-binding; DNA-binding;

KW Nuclear protein; Repeat.

FT DOMAIN 443 987 DOSAGE COMPENSATION DOMAIN 1.

FT DOMAIN 1508 1516 SEX DETERMINATION DOMAIN.

FT DOMAIN 2080 2105 DOSAGE COMPENSATION DOMAIN 2.

FT ZN_FING 2078 2105 C2H2-TYPE.

FT ZN_FING 2117 2141 C2H2-TYPE.

SQ SEQUENCE 2150 AA; 249954 MW; 7430D77AC784EA46 CRC64;

Query Match 60.0%; Score 33; DB 1; Length 2150;

Best Local Similarity 50.0%; Pred. No. 1.8e+02;

Matches 5; Conservative 2; Mismatches 3; Indels 0; Gaps 0;

OY 1 HHOKLVFAE 10

DB 2140 HHSRRCEFFAD 2149

RESULT 34

G6PI_KLEOX STANDARD; PRT; 167 AA.

ID G6PI_KLEOX

AC P77877;

DT 15-JUL-1998 (Rel. 36, Created)

DT 15-JUL-1998 (Rel. 36, Last sequence update)

DT 01-MAR-2002 (Rel. 41, Last annotation update)

DE Glucose-6-phosphate isomerase (EC 5.3.1.9) (GPI) (Phosphoglucose isomerase) (PGI) (Phosphohexose isomerase) (PHI) (Fragment).

GN PGI.

OS Klebsiella oxytoca.

OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;

OC Klebsiella.

OX NCBI_TaxID=571;

RN [1]

RP SEQUENCE FROM N.A.

RC STRAIN-EA321;

RX MEDLINE-97032593; PubMed=8875859;

RA Katz L.A.;

RT "Transkingdom transfer of the phosphoglucose isomerase gene.";

RL J. Mol. Evol. 43:453-459(1996).

CC -!- CATALYTIC ACTIVITY: D-glucose 6-phosphate -> D-fructose 6-phosphate.

CC -!- PATHWAY: INVOLVED IN GLYCOLYSIS AND IN GLUCONEOGENESIS.

CC -!- SUBCELLULAR LOCATION: Cytoplasmic.

CC -!- SIMILARITY: BELONGS TO THE GPI FAMILY.

CC -----

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CC -----

DR EMBL; U54763; AAB50058.1; -

DR InterPro; IPR001672; G6P_Isomerase.

DR Pfam; PF00342; PGI; 1.

DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; PARTIAL.

DR PROSITE; PS00174; P_GLUCOSE_ISOMERASE_2; PARTIAL.

KW Isomerase; Gluconeogenesis; Glycolysis.

FT NON_TER 167 167

SQ SEQUENCE 167 AA; 18875 MW; F6C56A969F06F891 CRC64;

Query Match 58.2%; Score 32; DB 1; Length 167;

Best Local Similarity 58.3%; Pred. No. 21;

Matches 7; Conservative 2; Mismatches 1; Indels 2; Gaps 1;

OY 1 HHOKLV--FFAE 10

DB 115 HHPKLLSNFFAQ 126

RESULT 35

CALC_MOUSE STANDARD; PRT; 178 AA.
AC Q63811;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Calcineurin B subunit isoform 2 (protein phosphatase 2B regulatory subunit 2) (protein phosphatase 3 regulatory subunit B alpha isoform 2).
GN PPP3R2.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
NCBI_TaxID=10090;
RN [1]
SEQUENCE FROM N.A.
RX MEDLINE=92392379; PubMed=1325794;
RA Ueki K., Muramatsu T., Kincaid R.L.;
RT "Structure and expression of two isoforms of the murine calmodulin-dependent protein phosphatase regulatory subunit (calcineurin B)."; Biochem. Biophys. Res. Commun. 187:537-543(1992).
RL Biochem. Biophys. Res. Commun. 187:537-543(1992).
CC -!- FUNCTION: REGULATORY SUBUNIT OF CALCINEURIN, A CALCIUM-DEPENDENT, CALMODULIN STIMULATED PROTEIN PHOSPHATASE. CONPERS CALCIUM SENSITIVITY.
CC -!- SUBUNIT: COMPOSED OF A CATALYTIC SUBUNIT (A) AND A REGULATORY SUBUNIT (B).
CC -!- TISSUE SPECIFICITY: TESTIS-SPECIFIC.
CC -!- MISCELLANEOUS: THIS PROTEIN HAS FOUR FUNCTIONAL CALCIUM-BINDING SITES.
CC -!- SIMILARITY: CONTAINS 4 EF-HAND CALCIUM-BINDING DOMAINS.
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CC -----
DR EMBL; S43865; AAB23172.1; -.
DR HSSP; P06705; LTCO.
DR MGD; MGI:107171; Ppp3r2.
DR InterPro; IPR002048; EF-hand.
DR InterPro; IPR001125; Recoverin.
DR Pfam; PF00036; ehand; 4.
DR PRINTS; PR00450; RECOVERIN.
DR SMART; SM00054; EFh; 4.
DR PROSITE; PS00018; EF_HAND; 4.
KW Calcium-binding; Repeat; Myristate.
KW INIT_MET 0
FT LIPID 0
FT CA_BIND 1 1 MYRISTATE (BY SIMILARITY).
FT CA_BIND 30 41 EF-HAND 1.
FT CA_BIND 62 73 EF-HAND 2.
FT CA_BIND 99 110 EF-HAND 3.
FT CA_BIND 140 151 EF-HAND 4.
SQ SEQUENCE 178 AA; 20528 MW; F453B9A047C240F5 CRC64;
Query Match 58.2%; Score 32; DB 1; Length 178;
Best Local Similarity 66.7%; Pred. No. 23;
Matches 6; Conservative 1; Mismatches 2; Indels 0; Gaps 0;
QY 2 HQKLVEFAE 10
|:|:| | |
Db 162 HKKLVEFE 170
RESULT 36
CYF_GUTH
ID CYF_GUTH STANDARD; PRT; 321 AA.
AC O78494;
DT 15-DEC-1998 (Rel. 37, Created)
DT 15-DEC-1998 (Rel. 37, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)

DE Apocytochrome F precursor.
GN PETA.
OS Guillardia theta (Cryptomonas phi).
OG Chloroplast.
OC Eukaryota; Cryptophyta; Cryptomonadaceae; Guillardia.
OX NCBI_TaxID=55529;
RN [1]
SEQUENCE FROM N.A.
RX MEDLINE=99128221; PubMed=9929392;
RA Douglas S.E., Penny S.L.;
RT "The plastid genome of the cryptophyte alga, Guillardia theta: complete sequence and conserved syntenic groups confirm its common ancestry with red algae."; J. Mol. Evol. 48:236-244(1999).
CC -!- FUNCTION: TRANSLOCATES PROTONS ACROSS THE THYLAKOID MEMBRANE AND TRANSFERS ELECTRONS FROM PHOTOSYSTEM II TO PHOTOSYSTEM I. IT RECEIVES ELECTRONS FROM THE RIESKE IRON-SULFUR PROTEIN AND PASSES THEM TO PLASTOCYANIN; THIS FUNCTION IS VERY SIMILAR TO THAT OF MITOCHONDRIAL CYTOCHROME C1.
CC -!- SUBUNIT: MEMBER OF THE CYTOCHROME B6/F COMPLEX INCLUDING CYTOCHROME B6, CYTOCHROME F AND PROBABLY AN IRON SULFUR PROTEIN.
CC -!- SUBCELLULAR LOCATION: Chloroplast thylakoid membrane (probable).
CC -!- SIMILARITY: BELONGS TO THE CYTOCHROME C FAMILY.
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CC -----
DR EMBL; AF041468; AAC35685.1; -.
DR HSSP; P36438; IHGZ.
DR InterPro; IPR002325; Apocyt_F.
DR InterPro; IPR000345; Cytc_heme_bind.
DR Pfam; PF01333; Apocytochrome_F; 1.
DR PRINTS; PR00610; CYTOCHROME_F.
DR PROSITE; PS00190; CYTOCHROME_C; 1.
KW Electron transport; Heme; Chloroplast; Thylakoid;
KW Photosynthesis; Photosystem I; Photosystem II; Transit peptide;
KW Transmembrane.
KW TRANSIT 1 38 CHLOROPLAST (BY SIMILARITY).
FT CHAIN 39 321 APOCYTOCHROME F.
FT BINDING 59 59 HEME (COVALENT) (PROBABLE).
FT BINDING 62 62 HEME (COVALENT) (PROBABLE).
FT METAL 63 63 IRON (HEME AXIAL LIGAND) (PROBABLE).
FT TRANSMEM 287 307 POTENTIAL.
SQ SEQUENCE 321 AA; 35173 MW; 42A1FF89FB05AE3D CRC64;
Query Match 58.2%; Score 32; DB 1; Length 321;
Best Local Similarity 100.0%; Pred. No. 42;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 2 HQKLVP 7
| | | | |
Db 161 HQKLVP 166
RESULT 37
FD3E_SOYBN
ID FD3E_SOYBN STANDARD; PRT; 380 AA.
AC P48625;
DT 01-FEB-1996 (Rel. 33, Created)
DT 01-FEB-1996 (Rel. 33, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Omega-3 fatty acid desaturase, endoplasmic reticulum (EC 1.14.99.-).
GN FAD3.
OS Glycine max (Soybean).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae; eurosids I; Fabales; Fabaceae; Papilionoideae; Phaseoleae; Glycine.
OX NCBI_TaxID=3847;

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RN RP SEQUENCE FROM N.A.
RC TISSUE=Seed:
RX MEDLINE=94302147; PubMed=8029334;
RA Kadev N.S., Wierzbicki A., Aegerter M., Caster C.S., Perez-Grau L.,
RA Kinney A.J., Hitz W.D., Booth J.R. Jr., Schweiger B., Stecca K.L.,
RA Allen S.M., Blackwell M., Reiter R.S., Carlson T.J., Russell S.H.,
RA Feldmann K.A., Pierce J., Browne J.;
RT "Cloning of higher plant omega-3 fatty acid desaturases.";
RL Plant Physiol. 103:467-476(1993).
CC
CC -1- FUNCTION: MICROSOMAL (ER) OMEGA-3 FATTY ACID DESATURASE INTRODUCES
CC THE THIRD DOUBLEBOND IN THE BIOSYNTHESIS OF 18:3 FATTY ACIDS,
CC IMPORTANT CONSTITUENTS OF PLANT MEMBRANES. IT IS THOUGHT TO USE
CC CYTOCHROME B5 AS AN ELECTRON DONOR AND TO ACT ON FATTY ACIDS
CC ESTERIFIED TO PHOSPHATIDYLCHOLINE AND, POSSIBLY, OTHER
CC PHOSPHOLIPIDS.
CC
CC -1- PATHWAY: POLYUNSATURATED FATTY ACID BIOSYNTHESIS.
CC -1- SUBCELLULAR LOCATION: Endoplasmic reticulum.
CC -1- DOMAIN: THE HISTIDINE BOX DOMAINS MAY CONTAIN THE ACTIVE SITE
CC AND/ OR BE INVOLVED IN METAL ION BINDING.
CC
CC -1- SIMILARITY: BELONGS TO THE FATTY ACID DESATURASE FAMILY.
CC
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CC
CC -----
CC EMBL; L22964; AAA61777.1; -.
CC
CC DR PIR; J02338; J02338.
CC
CC DR InterPro; IPR001225; FA_desaturase.
CC
CC DR Pfam; PF00487; FA_desaturase; 1.
CC
CC DR ProDom; PD001081; FA_desaturase; 1.
CC
CC KW Oxidoreductase; Fatty acid biosynthesis; Endoplasmic reticulum;
CC Transmembrane.
CC
CC FT TRANSMEM 55 75 POTENTIAL.
CC FT TRANSMEM 212 232 POTENTIAL.
CC FT TRANSMEM 236 256 POTENTIAL.
CC FT DOMAIN 100 104 HISTIDINE BOX 1.
CC FT DOMAIN 136 140 HISTIDINE BOX 2.
CC FT DOMAIN 303 307 HISTIDINE BOX 3.
CC
CC SQ SEQUENCE 380 AA; 44185 MW; BF800F93CF4C29D7 CRC64;

Query Match 58.28; Score 32; DB 1; Length 380;
Best Local Similarity 62.58; Pred. No. 50;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Oy 1 HHQKLVFF 8
    ||||| ::
Db 264 HHQKLPWY 271

RESULT 38
LBP_HUMAN STANDARD; PRT: 481 AA.
AC P18428; Q92672; Q43438; Q9UD66; Q9H403;
DT 01-NOV-1990 (Rel. 16, Created)
DT 15-DEC-1998 (Rel. 37, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Lipopolysaccharide-binding protein precursor (LBP).
GN LBP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
[1]
RN RP SEQUENCE FROM N.A.
RX MEDLINE=90385281; PubMed=2402637;
RA Schumann R.R., Leong S.R., Flagg G.W., Gray P.W., Wright S.D.,
RA Mathison J.C., Tobias P.S., Ulevitch R.J.;
RT "Structure and function of lipopolysaccharide binding protein.";
```

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RL Science 249:1429-1431(1990).
[2]
RN RP SEQUENCE FROM N.A.
RX MEDLINE=94292492; PubMed=7517398;
RA Wilde C.G., Seilhamer J.J., McGrogan M., Ashton N., Snable J.L.,
RA Lane J.C., Leong S.R., Thornton M.B., Miller K.L., Scott R.W.;
RT "Bactericidal/permeability-increasing protein and lipopolysaccharide
RT (LPS)-binding protein: LPS binding properties and effects on LPS-
RT mediated cell activation.";
RL J. Biol. Chem. 269:17411-17416(1994).
[3]
RN RP SEQUENCE FROM N.A.
RX Hubacek J.A., Aslanidis C., Schmitz G.;
RL Submitted (OCT-1996) to the EMBL/GenBank/DBJ databases.
[4]
RN RP SEQUENCE FROM N.A.
RX MEDLINE=98110577; PubMed=9441745;
RA Kirschning C.J., Au-Young J., Lamping N., Reuter D., Pfeil D.,
RA Seilhamer J.J., Schumann R.R.;
RT "Similar organization of the lipopolysaccharide-binding protein (LBP)
RT and phospholipid transfer protein (PLTP) genes suggests a common gene
RT family of lipid-binding proteins.";
RL Genomics 46:416-425(1997).
[5]
RN RP SEQUENCE FROM N.A.
RX TISSUE=Liver;
RL Long J.Y., Liu J.Q., Xue Y.N., Wang H.X.;
RT "Cloning and sequencing of human lipopolysaccharide-binding protein
RT gene.";
RL Sheng Wu Huaxue Yu Shengwu Wuli Jinzhan 25:469-471(1998).
[6]
RN RP SEQUENCE FROM N.A.
RX Deloukas P., Matthews L.H., Ashurst J., Burton J., Gilbert J.G.R.,
RA Jones M., Stavrides G., Almeida J.P., Babbage A.K., Bagguely C.L.,
RA Bailey J., Barlow K.F., Bates K.N., Beard L.M., Beare D.M.,
RA Beasley O.P., Bird C.P., Blakey S.E., Bridgeman A.M., Brown A.J.,
RA Buck D., Burrill W., Butler A.P., Carder C., Carter N.P.,
RA Chapman J.C., Clamp M., Clark G., Clark L.N., Clark S.V., Clee C.M.,
RA Clegg S., Cobley V.E., Collier R.E., Connor R., Corby N.R.,
RA Coulson A., Coville G.J., Deadman R., Dhami P., Dunn M.,
RA Ellington A.G., Frankland J.A., Fraser A., French L., Garner P.,
RA Grafham D.V., Griffiths C., Griffiths M.N.D., Gwilliam R., Hall R.E.,
RA Hammond S., Harley J.L., Heath P.D., Ho S., Holden J.L., Howden P.J.,
RA Huckle E., Hunt A.R., Hunt S.E., Jekosch K., Johnson C.M., Johnson D.,
RA Kay M.P., Kimberley A.M., King A., Knights A., Laird G.K., Lawlor S.,
RA Leivasalho M.H., Leversha M., Lloyd D.M., Lloyd D.M., Lovell J.D.,
RA Marsh V.L., Martin S.L., McConachie L.J., McElay K., McMurray A.A.,
RA Milne S., Mistry D., Moore M.J.F., Mullikin J.C., Nickerson T.,
RA Oliver K., Parker A., Patel R., Pearce T.A.V., Peck A.I.,
RA Phillimore B.J.C.T., Prathalingam S.R., Plumb R.W., Ramsay H.,
RA Rice C.M., Ross M.T., Scott C.E., Sehra H.K., Shownkeen R., Sims S.,
RA Skuce C.D., Smith M.L., Soderlund C., Steward C.A., Sulston J.E.,
RA Swann M., Sycamore N., Taylor R., Tee L., Thomas D.W., Thorpe A.,
RA Tracey A., Tromans A.C., Vaudin M., Wall M., Wallis J.M.,
RA Whitehead S.I., Whittaker P., Willey D.L., Williams L., Williams S.A.,
RA Wilming L., Wray P.W., Hubbard T., Durbin R.M., Bentley D.R., Beck S.,
RA Rogers J.;
RT "The DNA sequence and comparative analysis of human chromosome 20.";
RL Nature 414:865-871(2001).
[7]
RN RP SEQUENCE OF 1-41 FROM N.A.
RX Sutton C.L., Smith R.I.F., Centola M.B., Theofan G.;
RL Submitted (MAY-1995) to the EMBL/GenBank/DBJ databases.
[8]
RN RP 3D-STRUCTURE MODELING.
RX MEDLINE=98227852; PubMed=9568897;
RA Beamer L.J., Carroll S.F., Eisenberg D.;
RT "The BPI/LBP family of proteins: a structural analysis of conserved
RT regions.";
RL Protein Sci. 7:906-914(1998).
CC -1- FUNCTION: BINDS TO THE LIPID A MOIETY OF BACTERIAL
CC LIPOPOLYSACCHARIDES (LPS), A GLYCOLIPID PRESENT IN THE OUTER
CC MEMBRANE OF ALL GRAM-NEGATIVE BACTERIA. THE LBP/LPS COMPLEX SEEMS
```


DT 30-MAY-2000 (Rel. 39, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE UDP-N-acetylmuramate--alanine ligase (EC 6.3.2.8) (UDP-N-
 DE acetylmuramoyl-L-alanine synthetase).
 GN MURC OR RP247.
 OS Rickettsia prowazekii.
 OC Bacteria; Proteobacteria; alpha subdivision; Rickettsiales;
 OC Rickettsiaceae; Rickettsiae; Rickettsia.
 OX NCBI_TaxID=782;
 RN [1]
 RP SEQUENCE FROM N.A.
 RC STRAIN=MADRID E;
 RX MEDLINE=99039439; PubMed=9823893;
 RA Andersson S.G.E., Zomorodipour A., Andersson J.O.,
 RA Sacheritz-Ponten T., Alsmark U.C.M., Podowski R.M., Naeslund A.K.,
 RA Eriksson A.-S., Winkler H.H., Kurland C.G.;
 RT "The genome sequence of Rickettsia prowazekii and the origin of
 RT mitochondria";
 RN Nature 396:133-140(1998).
 RL CC -1- FUNCTION: CELL WALL FORMATION (BY SIMILARITY).
 CC -1- CATALYTIC ACTIVITY: ATP + UDP-N-acetylmuramoyl + L-alanine = ADP +
 CC phosphate + UDP-N-acetylmuramoyl-L-alanine.
 CC -1- PATHWAY: PEPTIDOGLYCAN BIOSYNTHESIS.
 CC -1- SUBCELLULAR LOCATION: Cytoplasmic (Probable).
 CC -1- SIMILARITY: BELONGS TO THE MURCDEF FAMILY.
 CC -----
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 CC -----
 DR EMBL: AJ235271; CAAL4709.1; -
 DR InterPro: IPR000713; Mur_ligase.
 DR Pfam: PF01225; Mur_ligase.C.
 DR Pfam: PF02875; Mur_ligase.C; 1.
 KW Peptidoglycan synthesis; Cell wall; Cell division; Ligase;
 KW ATP-binding; Complete proteome.
 FT NP_BIND 120 126 ATP (POTENTIAL).
 SQ SEQUENCE 495 AA; 54612 MW; 2E18464088FAD2D6 CRC64;

 Query Match 58.2%; Score 32; DB 1; Length 495;
 Best Local Similarity 60.0%; Pred. No. 65;
 Matches 6; Conservative 0; Mismatches 4; Indels 0; Gaps 0;

 QY 1 HHKQLVFAE 10
 ||| |||
 Db 428 HHDKANFLAE 437

 RESULT 41
 ACHB_MOUSE ID ACHB_MOUSE STANDARD; PRT; 501 AA.
 AC P09690;
 DT 01-MAR-1989 (Rel. 10, Created)
 DT 01-MAR-1989 (Rel. 10, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Acetylcholine receptor protein, beta chain precursor.
 GN CHRN1 OR ACRB.
 OS Mus musculus (Mouse).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 OX NCBI_TaxID=10090;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=87057335; PubMed=3782129;
 RA Buonanno A., Mudd J., Shah V., Merlie J.P.;
 RT "A universal oligonucleotide probe for acetylcholine receptor genes.
 RT Selection and sequencing of cDNA clones for the mouse muscle beta
 RT subunit.";

RL J. Biol. Chem. 261:16451-16458(1986).
 RN [2]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=89214211; PubMed=2708381;
 RA Buonanno A., Mudd J., Merlie J.P.;
 RT "Isolation and characterization of the beta and epsilon subunit genes
 RT of mouse muscle acetylcholine receptor.";
 RL J. Biol. Chem. 264:7611-7616(1989).
 CC -1- FUNCTION: AFTER BINDING ACETYLCHOLINE, THE ACHR RESPONDS BY AN
 CC EXTENSIVE CHANGE IN CONFORMATION THAT AFFECTS ALL SUBUNITS AND
 CC LEADS TO OPENING OF AN ION-CONDUCTING CHANNEL ACROSS THE PLASMA
 CC MEMBRANE.
 CC -1- SUBUNIT: PENTAMER OF TWO ALPHA CHAINS, AND ONE EACH OF THE BETA,
 CC DELTA, AND GAMMA (IN IMMATURE MUSCLE) OR EPSILON (IN MATURE
 CC MUSCLE) CHAINS.
 CC -1- SUBCELLULAR LOCATION: Integral membrane protein.
 CC -1- SIMILARITY: BELONGS TO THE LIGAND-GATED IONIC CHANNELS FAMILY.
 CC -----
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 CC or send an email to license@isb-sib.ch).
 CC -----
 DR EMBL: M14537; AAA37154.1; -
 DR EMBL: J04699; AAA37156.1; -
 DR PIR: A25338; A25338.
 DR MGD; MGI:87890; Chrnbl.
 DR InterPro: IPR000188; GABAA_receptor.
 DR InterPro: IPR001175; Neur_channel.
 DR Pfam: PF02931; Neur_chan_LBD; 1.
 DR Pfam: PF02932; Neur_chan_memb; 1.
 DR PRINTS; PR00252; NRIONCHANNEL.
 DR PROSITE; PS00236; NEUOTR_ION_CHANNEL; 1.
 KW Receptor; Postsynaptic membrane; Ionic channel; Glycoprotein; Signal;
 KW Transmembrane; Phosphorylation.
 FT SIGNAL 1 23
 FT CHAIN 24 501 ACETYLCHOLINE RECEPTOR PROTEIN, BETA
 FT CHAIN EXTRACELLULAR.
 FT DOMAIN 24 244
 FT TRANSMEM 245 269
 FT TRANSMEM 277 295
 FT TRANSMEM 311 332
 FT DOMAIN 333 469 CYTOPLASMIC.
 FT TRANSMEM 470 488
 FT DISULFID 151 165 BY SIMILARITY.
 FT CARBOHYD 164 164 N-LINKED (GLCNAC...) (PROBABLE).
 FT MOD_RES 390 390 PHOSPHORYLATION (BY TYR-KINASES)
 FT (BY SIMILARITY).
 SQ SEQUENCE 501 AA; 56930 MW; 787BDDA90EBB0EF2 CRC64;

 Query Match 58.2%; Score 32; DB 1; Length 501;
 Best Local Similarity 37.5%; Pred. No. 66;
 Matches 3; Conservative 5; Mismatches 0; Indels 0; Gaps 0;

 QY 1 HHKLVFF 8
 ||:::|:
 Db 231 HHEEVIFY 238

 RESULT 42
 ACHG_HUMAN ID ACHG_HUMAN STANDARD; PRT; 517 AA.
 AC P07510;
 DT 01-APR-1988 (Rel. 07, Created)
 DT 01-APR-1988 (Rel. 07, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Acetylcholine receptor protein, gamma chain precursor.
 GN CHRG OR ACHRG.
 OS Homo sapiens (Human).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

OC Mammalia; Eutheria; Primates; Catarrhini; Hominoidea; Homo.
 OX NCBI_TaxID=9606;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=85101368; PubMed=3967651;
 RA Shibahara S., Kubo T., Ferski H.J., Takahashi H., Noda M., Numa S.;
 RT "Cloning and sequence analysis of human genomic DNA encoding gamma
 subunit precursor of muscle acetylcholine receptor.";
 RL Eur. J. Biochem. 146:15-22(1985).
 RN [2]
 RP SEQUENCE FROM N.A.
 RX TISSUE=Muscle fibroblast;
 RA MEDLINE=93345508; PubMed=768301;
 RA Beeson D.M.W., Brydson M., Betty M., Jeremiah S., Povey S.,
 RA Vincent A., Newson-Davis J.;
 RT "Primary structure of the human muscle acetylcholine receptor. cDNA
 cloning of the gamma and epsilon subunits.";
 RL Eur. J. Biochem. 215:229-238(1993).
 CC -!- FUNCTION: AFTER BINDING ACETYLCHOLINE, THE ACHR RESPONDS BY AN
 CC EXTENSIVE CHANGE IN CONFORMATION THAT AFFECTS ALL SUBUNITS AND
 CC LEADS TO OPENING OF AN ION-CONDUCTING CHANNEL ACROSS THE PLASMA
 CC MEMBRANE.
 CC -!- SUBUNIT: PENTAMER OF TWO ALPHA CHAINS, AND ONE EACH OF THE BETA,
 CC DELTA, AND GAMMA (IN IMMATURE MUSCLE) OR EPSILON (IN MATURE
 CC MUSCLE) CHAINS.
 CC -!- SUBCELLULAR LOCATION: Integral membrane protein.
 CC -!- SIMILARITY: BELONGS TO THE LIGAND-GATED IONIC CHANNELS FAMILY.
 CC
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 CC
 DR EMBL; M1811; AA51568.2; JOINED.
 DR EMBL; L29197; AA51568.2; JOINED.
 DR EMBL; X01715; CAA25861.1; ALT SEQ.
 DR EMBL; X01716; CAA25861.1; JOINED.
 DR EMBL; X01717; CAA25861.1; JOINED.
 DR EMBL; X01718; CAA25861.1; JOINED.
 DR EMBL; X01719; CAA25861.1; JOINED.
 DR EMBL; X01720; CAA25861.1; JOINED.
 DR EMBL; X01721; CAA25861.1; JOINED.
 DR EMBL; X04759; CAA25861.1; JOINED.
 DR PIR; A23261; A23261.
 DR PIR; S34776; S34776.
 DR MIM; 100730; -
 DR InterPro; IPR000188; GABAA_receptor.
 DR InterPro; IPR001175; Neur_channel.
 DR Pfam; PF02931; Neur_chan_LBD; 1.
 DR Pfam; PF02932; Neur_chan_memb; 1.
 DR PRINTS; PR00252; NRIONCHANNEL.
 DR PROSITE; PS00236; NEUROTR_ION_CHANNEL; 1.
 DR Receptor; Postsynaptic membrane; Ionic channel; Glycoprotein; Signal;
 KW Transmembrane.
 FT SIGNAL 1 22
 FT CHAIN 23 517
 FT
 FT DOMAIN 23 240
 FT TRANSMEM 241 265
 FT TRANSMEM 275 293
 FT TRANSMEM 309 330
 FT DOMAIN 331 474
 FT TRANSMEM 475 495
 FT TRANSMEM 150 164
 FT DISULFID 52 52
 FT CARBOHYD 163 163
 FT CARBOHYD 517 AA; 57897 MW; D4587257087E102C CRC64;
 SQ SEQUENCE 517 AA; 57897 MW; D4587257087E102C CRC64;
 Query Match 58.2%; Score 32; DB 1; Length 517;
 Best Local Similarity 71.4%; Pred. No. 68;
 Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;
 QY 2 HOKLVFF 8
 DB 228 HOKVVFY 234
 RESULT 43
 ACHG_BOVIN
 ID ACHG_BOVIN STANDARD; PRT; 519 AA.
 AC P13536;
 DT 01-JAN-1990 (Rel. 13; Created)
 DT 01-JAN-1990 (Rel. 13; Last sequence update)
 DT 01-OCT-1996 (Rel. 34; Last annotation update)
 DE Acetylcholine receptor protein, gamma chain precursor.
 GN CHNG.
 OS Bos taurus (Bovine).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae;
 OC Bovidae; Bovinae; Bos.
 OX NCBI_TaxID=9913;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=84285374; PubMed=6547904;
 RA Takai T., Noda M., Furutani Y., Takahashi H., Notake M., Shimizu S.,
 RA Kavano T., Tanabe T., Tanaka K.-I., Hirose T., Inayama S., Numa S.;
 RT "Primary structure of gamma subunit precursor of calf-muscle
 acetylcholine receptor deduced from the cDNA sequence.";
 RL Eur. J. Biochem. 143:109-115(1984).
 CC -!- FUNCTION: AFTER BINDING ACETYLCHOLINE, THE ACHR RESPONDS BY AN
 CC EXTENSIVE CHANGE IN CONFORMATION THAT AFFECTS ALL SUBUNITS AND
 CC LEADS TO OPENING OF AN ION-CONDUCTING CHANNEL ACROSS THE PLASMA
 CC MEMBRANE.
 CC -!- SUBUNIT: PENTAMER OF TWO ALPHA CHAINS, AND ONE EACH OF THE BETA,
 CC DELTA, AND GAMMA (IN IMMATURE MUSCLE) OR EPSILON (IN MATURE
 CC MUSCLE) CHAINS.
 CC -!- SUBCELLULAR LOCATION: Integral membrane protein.
 CC -!- SIMILARITY: BELONGS TO THE LIGAND-GATED IONIC CHANNELS FAMILY.
 CC
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 CC
 DR EMBL; M28307; AAA30351.1; -
 DR InterPro; IPR000188; GABAA_receptor.
 DR InterPro; IPR001175; Neur_channel.
 DR Pfam; PF02931; Neur_chan_LBD; 1.
 DR Pfam; PF02932; Neur_chan_memb; 1.
 DR PRINTS; PR00252; NRIONCHANNEL.
 DR PROSITE; PS00236; NEUROTR_ION_CHANNEL; 1.
 DR Receptor; Postsynaptic membrane; Ionic channel; Glycoprotein; Signal;
 KW Transmembrane.
 FT SIGNAL 1 22
 FT CHAIN 23 519
 FT
 FT DOMAIN 23 240
 FT TRANSMEM 241 265
 FT TRANSMEM 274 292
 FT TRANSMEM 308 329
 FT DOMAIN 330 476
 FT TRANSMEM 477 497
 FT TRANSMEM 150 164
 FT DISULFID 52 52
 FT CARBOHYD 163 163
 FT CARBOHYD 519 AA; 58178 MW; B72DE5487F7B5C4E CRC64;
 SQ SEQUENCE 519 AA; 58178 MW; B72DE5487F7B5C4E CRC64;
 Query Match 58.2%; Score 32; DB 1; Length 519;
 Best Local Similarity 71.4%; Pred. No. 68;
 Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

RX MEDLINE=20013071; PubMed=10543953;
RA Weyand M., Hecht H.-J., Kiess M., Liaud M.-F., Vilter H.,
RA Schomburg D.;
RT "X-ray structure determination of a vanadium-dependent
RT haloperoxidase from Ascophyllium nodosum at 2.0-A resolution.";
RL J. Mol. Biol. 293:595-611(1999).
CC [2]
CC SEQUENCE OF 320-556 FROM N.A., SEQUENCE OF 326-341; 383-426; 471-479
RP AND 481-556, AND FUNCTION.
RX MEDLINE=96081028; PubMed=8564812;
RA Vilter H.;
RT "Vanadium-dependent haloperoxidases";
RL (In) Sigel H., Sigel A. (eds.);
RL Metal ions in biological system-vanadium and its role in life,
RL pp. 31-325-362 Marcel Dekker, New York (1995).
CC -!- CATALYTIC ACTIVITY: Halide + H(2)O(2) + H(+) = HOHAL + H(2)O.
CC -!- COFACTOR: VANADIUM.
CC -!- SUBUNIT: HOMODIMER LINKED BY TWO INTERCHAIN DISULFIDE BONDS.
CC -!- SIMILARITY: TO OTHER BACTERIAL NON-HEME BROMO- AND CHLORO-
CC PEROXIDASES.
DR PDB: 1OI9; 10-JUN-00.
DR InterPro: IPR000326; PA_PTPase.
KW Oxidoreductase; Peroxidase; Vanadium; 3D-structure.
FT MOD_RES 1 1 PYRROLIDONE CARBOXYLIC ACID.
FT DISULFID 3 3 INTERCHAIN (WITH C-41 OF OTHER CHAIN).
FT DISULFID 41 41 INTERCHAIN (WITH C-3 OF OTHER CHAIN).
FT DISULFID 77 86
FT DISULFID 441 462
FT DISULFID 544 555
FT ACT_SITE 411 411
FT ACT_SITE 418 418
FT METAL 486 486 VANADIUM.
FT CONFLICT 321 321 S -> D (IN REF. 2).
FT CONFLICT 341 341 K -> N (IN REF. 2; DNA SEQUENCE).
FT CONFLICT 403 404 AI -> VY (IN REF. 2; DNA SEQUENCE).
FT CONFLICT 407 408 GS -> T (IN REF. 2).
FT CONFLICT 409 409 P -> S (IN REF. 2; AA SEQUENCE).
FT CONFLICT 441 444 CYPD -> AIR (IN REF. 2).
FT CONFLICT 470 470 N -> K (IN REF. 2).
SQ SEQUENCE 557 AA; 60343 MW; E3D8557AB92B16F4 CRC64;
Query Match 58.2%; Score 32; DB 1; Length 557;
Best Local Similarity 66.7%; Pred. No. 73;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;
QY 2 HOKLVFFAE 10
Db 507 HQELMTFAE 515
|||:|:|
|||:|:|
RESULT 47
YAWG_SCHPO
ID YAWG_SCHPO STANDARD; PRT; 616 AA.
AC Q10190;
DT 01-OCT-1996 (Rel. 34, Created)
DT 01-OCT-1996 (Rel. 34, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Hypothetical GTP-binding protein C3F10.16C in chromosome I.
GN SPAC3F10.16C.
OS Schizosaccharomyces pombe (Fission yeast).
OC Eukaryota; Fungi; Ascomycota; Schizosaccharomycetes;
OC Schizosaccharomycetales; Schizosaccharomycetaceae;
OC Schizosaccharomycetes.
OX NCBI_TaxID=4896;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=972.
RA Murphy L., Harris D., Barrell B.G., Rajandream M.A., Walsh S.V.;
RL Submitted (FEB-1996) to the EMBL/GenBank/DBJ databases.
CC -!- SIMILARITY: BELONGS TO THE MMRL/HSR1 FAMILY OF GTP-BINDING
CC PROTEINS.
CC -----
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RP SEQUENCE OF 203-306 FROM N.A.
RX MEDLINE=88030021; PubMed=3666131;
RA Witzemann V., Barg B., Nishikawa Y., Sakmann B., Numa S.;
RT "Differential regulation of muscle acetylcholine receptor gamma- and
RT epsilon-subunit mRNAs";
RL FEBS Lett. 223:104-112(1987).
CC -!- FUNCTION: AFTER BINDING ACETYLCHOLINE, THE ACHR RESPONDS BY AN
CC EXTENSIVE CHANGE IN CONFORMATION THAT AFFECTS ALL SUBUNITS AND
CC LEADS TO OPENING OF AN ION-CONDUCTING CHANNEL ACROSS THE PLASMA
CC MEMBRANE.
CC -!- SUBUNIT: PENTAMER OF TWO ALPHA CHAINS, AND ONE EACH OF THE BETA,
CC DELTA, AND GAMMA (IN IMMATURE MUSCLE) OR EPSILON (IN MATURE
CC MUSCLE) CHAINS.
CC -!- SUBCELLULAR LOCATION: Integral membrane protein.
CC -!- SIMILARITY: BELONGS TO THE LIGAND-GATED IONIC CHANNELS FAMILY.
CC -----
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CC -----
DR EMBL: X74834; CAA52828.1; -;
DR EMBL: X66364; CAA29662.1; -;
DR PIR: S03082; S03082.
DR PIR: S13874; S13874.
DR InterPro: IPR000188; GABAA_receptor.
DR InterPro: IPR001175; Neur_chan.
DR Pfam: PF02931; Neur_chan_LBD; 1.
DR Pfam: PF02932; Neur_chan_memb; 1.
DR PRINTS: PR00252; NRIONCHANNEL.
DR PROSITE: PS00236; NEUROTR_ION_CHANNEL; 1.
KW Receptor; Postsynaptic membrane; Ionic channel; Glycoprotein; Signal;
KW Transmembrane.
FT SIGNAL 1 22 BY SIMILARITY.
FT CHAIN 23 519 ACETYLCHOLINE RECEPTOR PROTEIN, GAMMA
FT CHAIN 23 519 CHAIN.
FT DOMAIN 23 240 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 241 265 POTENTIAL.
FT TRANSMEM 274 292 POTENTIAL.
FT TRANSMEM 308 329 POTENTIAL.
FT DOMAIN 330 476 CYTOPLASMIC (POTENTIAL).
FT TRANSMEM 477 497 POTENTIAL.
FT DISULFID 150 164 BY SIMILARITY.
FT CARBOHYD 52 52 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 163 163 N-LINKED (GLCNAC. . .) (PROBABLE).
SQ SEQUENCE 519 AA; 58621 MW; 1C97A83DE42A0D09 CRC64;
Query Match 58.2%; Score 32; DB 1; Length 519;
Best Local Similarity 71.4%; Pred. No. 68;
Matches 5; Conservative 2; Mismatches 0; Indels 0; Gaps 0;
QY 2 HOKLVFF 8
Db 228 HQKVVEY 234
|||:|:|
|||:|:|
RESULT 46
PRXV_ASCNO
ID PRXV_ASCNO STANDARD; PRT; 557 AA.
AC P81701;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Vanadium haloperoxidase (EC 1.11.1.-) (V-BPO).
OS Ascophyllium nodosum.
OC Eukaryota; stramenopiles; Phaeophyceae; Fucales; Fucaeae;
OC Ascophyllium.
OX NCBI_TaxID=52969;
RN [1]
RP SEQUENCE, X-RAY CRYSTALLOGRAPHY (2.0 ANGSTROMS), AND FUNCTION.

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CC -----

DR EMBL; Z69369; CAA93314.1; -
DR InterPro; IPR002917; MMR_HSR1.
DR Pfam; PF01926; MMR_HSR1; 1.
KW Hypothetical protein; GTP-binding.
FT NP_BIND 308 315 GTP (POTENTIAL).
FT NP_BIND 352 356 GTP (POTENTIAL).
SQ SEQUENCE 616 AA; 69674 MW; F02A2996AF06FB68 CRC64;

Query Match 58.2%; Score 32; DB 1; Length 616;
Best Local Similarity 83.3%; Pred. No. 81;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
|||||
Db 480 HHQKIV 485

RESULT 48
GLGB_AGRTU STANDARD; PRT; 734 AA.
AC P52979;
DT 01-OCT-1996 (Rel. 34, Created)
DT 15-DEC-1998 (Rel. 37, Last sequence update)
DT 15-JUL-1999 (Rel. 38, Last annotation update)
DE 1.4-alpha-glucan branching enzyme (EC 2.4.1.18) (Glycogen branching
DE enzyme).
GN GLGB.
OS Agrobacterium tumefaciens.
OC Bacteria; Proteobacteria; alpha subdivision; Rhizobiaceae group;
OC Rhizobiaceae; Rhizobium.
OX NCBI_TaxID=358;
RN [1]
RC SEQUENCE FROM N.A.
RP STRAIN-A348.
RX MEDLINE=99069330; PubMed=9851999;
RA Ugalde J.E., Lepek V., Uttaro A.D., Estrella J., Iglesias A.,
RA Ugalde R.A.;
RT "Gene organization and transcription analysis of the Agrobacterium
RT tumefaciens glycogen (glg) operon: two transcripts for the single
RT phosphoglucomutase gene.;
RL J. Bacteriol. 180:6557-6564(1998).
CC -1- CATALYTIC ACTIVITY: Formation of 1,6-glucosidic linkages of
CC glycogen.
CC -1- PATHWAY: THIRD STEP IN GLYCOGEN BIOSYNTHESIS.
CC -1- SUBUNIT: MONOMER (BY SIMILARITY).
CC -1- SIMILARITY: BELONGS TO FAMILY 13 OF GLYCOSYL HYDROLASES, ALSO
CC KNOWN AS THE ALPHA-AMYLASE FAMILY.
CC -----

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CC -----

DR EMBL; AF033856; AAD03472.1; -
DR InterPro; IPR000461; Alpha_amylase.
DR InterPro; IPR004193; Isoamylase_N.
DR Pfam; PF00128; alpha-amylase; 1.
DR Pfam; PF02922; isoamylase_N; 1.
KW Glycogen biosynthesis; Transferase; Glycosyltransferase.
FT ACT_SITE 417 417 BY SIMILARITY.
FT ACT_SITE 470 470 BY SIMILARITY.
FT ACT_SITE 538 538 BY SIMILARITY.
SQ SEQUENCE 734 AA; 83623 MW; 70A3CD35A77F31E6 CRC64;

Query Match 58.2%; Score 32; DB 1; Length 734;
Best Local Similarity 71.4%; Pred. No. 97;
Matches 5; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLV 7
|||||
Db 515 HHQELTF 521

RESULT 49
PLD_PIMBR STANDARD; PRT; 808 AA.
AC O04883;
DT 15-DEC-1998 (Rel. 37, Created)
DT 15-DEC-1998 (Rel. 37, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Phospholipase D precursor (EC 3.1.4.4) (PLD) (Choline phosphatase)
DE (Phosphatidylcholine-hydrolyzing phospholipase D).
GN PLD.
OS Pimpinella brachycarpa.
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC Asteridae; euasterids II; Apiales; Apiaceae; Pimpinella.
OX NCBI_TaxID=45043;
RN [1]
RP SEQUENCE FROM N.A.
RA Cha Y.-Y., Lee K.-W., Kim J.C., Han T.J., Lee W.S., Cho S.H.;
RT "Nucleotide sequence of a cDNA encoding phospholipase D from
RT Pimpinella brachycarpa.";
RL (In) Plant Gene Register PGR97-092.
CC -1- FUNCTION: PLAYS AN IMPORTANT ROLE IN VARIOUS CELLULAR PROCESSES.
CC -1- CATALYTIC ACTIVITY: A phosphatidylcholine + H(2)O -> choline + a
CC phosphatidate.
CC -1- COFACTOR: CALCIUM (BY SIMILARITY).
CC -1- SIMILARITY: BELONGS TO THE PHOSPHOLIPASE D FAMILY.
CC -1- SIMILARITY: CONTAINS 1 C2 DOMAIN.
CC -1- SIMILARITY: CONTAINS 2 PLDC DOMAINS.
CC -----

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CC -----

DR EMBL; U96438; AAB70463.1; -
DR InterPro; IPR000008; C2.
DR InterPro; IPR001736; PLD.
DR Pfam; PF00168; C2; 1.
DR Pfam; PF00614; PLDC; 2.
DR SMART; SM00239; C2; 1.
DR SMART; SM00155; PLDC; 2.
DR PROSITE; PS00004; C2_DOMAIN_2; 1.
KW Hydrolase; Lipid degradation; Calcium; Repeat.
FT PROPEP 1 2 POTENTIAL.
FT CHAIN 7 808 PHOSPHOLIPASE D.
FT DOMAIN 1 109 C2 DOMAIN.
FT DOMAIN 326 364 PLDC 1.
FT DOMAIN 654 681 PLDC 2.
SQ SEQUENCE 808 AA; 91672 MW; E83DA015B06F2164 CRC64;

Query Match 58.2%; Score 32; DB 1; Length 808;
Best Local Similarity 83.3%; Pred. No. 1.le+02;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
|||||
Db 330 HHQKIV 335

RESULT 50

Search completed: October 29, 2002, 09:24:48
Job time : 15 secs

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PLD_RICCO
ID   PLD_RICCO      STANDARD;          PRT;          808 AA.
AC   Q41142; P93507;
DT   01-NOV-1997 (Rel. 35, Created)
DT   01-NOV-1997 (Rel. 35, Last sequence update)
DT   16-OCT-2001 (Rel. 40, Last annotation update)
DE   Phospholipase D precursor (EC 3.1.4.4) (PLD) (Choline phosphatase)
DE   (Phosphatidylcholine-hydrolyzing phospholipase D).
OS   Ricinus communis (Castor bean).
OC   Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC   Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
OC   eurosids I; Malpighiales; Euphorbiaceae; Ricinus.
OX   NCBI_TaxID=3988;
RN   [1]
RP   SEQUENCE FROM N.A., AND PARTIAL SEQUENCE.
RX   STRAIN=CV. HALE; TISSUE=Endosperm;
RX   MEDLINE=94327597; PubMed=8051126;
RA   Wang X., Xu L., Zheng L.;
RT   "Cloning and expression of phosphatidylcholine-hydrolyzing
RT   phospholipase D from Ricinus communis L.";
RL   J. Biol. Chem. 269:20312-20317(1994).
RN   [2]
RP   SEQUENCE FROM N.A.
RC   TISSUE=Leaf;
RX   MEDLINE=97134969; PubMed=8980529;
RA   Xu L., Zheng L., Coughlan S.J., Wang X.;
RT   "Structure and analysis of phospholipase D gene from Ricinus communis
RT   L.";
RL   Plant Mol. Biol. 32:767-771(1996).
CC   -!- FUNCTION: PLAYS AN IMPORTANT ROLE IN CELLULAR PATHWAYS INCLUDING
CC   -!- SIGNAL TRANSDUCTION PATHWAYS.
CC   -!- CATALYTIC ACTIVITY: A phosphatidylcholine + H(2)O = choline + a
CC   phosphatidate.
CC   -!- COFACTOR: CALCIUM (BY SIMILARITY).
CC   -!- TISSUE SPECIFICITY: MOSTLY EXPRESSED IN VACUOLES, ENDOPLASMIC
CC   RETICULUM, A FEW IN PLASTIDS AND PLASMA MEMBRANE. EXPRESSION IS
CC   HIGHER IN RADICLE THAN IN ENDOSPERM.
CC   -!- SIMILARITY: BELONGS TO THE PHOSPHOLIPASE D FAMILY.
CC   -!- SIMILARITY: CONTAINS 1 C2 DOMAIN.
CC   -!- SIMILARITY: CONTAINS 2 PLDC DOMAINS.
CC   -----
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CC   or send an email to license@isb-sib.ch).
CC   -----
DR   EMBL; L33686; AAB04095.1; -.
DR   EMBL; U72693; AAB37305.1; -.
DR   InterPro; IPR000008; C2.
DR   InterPro; IPR001736; PLD.
DR   Pfam; PF00614; PLDC; 2.
DR   SMART; SM00239; C2; 1.
DR   SMART; SM00155; PLDC; 2.
DR   PROSITE; PS50004; C2_DOMAIN_2; 1.
KW   Hydrolase; Lipid degradation; Calcium; Repeat.
FT   PROPEP      1      30
FT   CHAIN       31      808  PHOSPHOLIPASE D.
FT   DOMAIN      1      109    C2 DOMAIN.
FT   DOMAIN      326    364    PLDC 1.
FT   DOMAIN      654    681    PLDC 2.
FT   CONFLICT    268    268    L -> I (IN REF. 2).
SQ   SEQUENCE 808 AA; 91992 MW; E75F6CFFB9ADF3CB CRC64;

Query Match      58.2%; Score 32; DB 1; Length 808;
Best Local Similarity 83.3%; Pred. No. 1.1e+02;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLV 6
    |||||
DB 330 HHQKIV 335
```

GenCore version 5.1.3
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OM protein - protein search, using sw model
Run on: October 29, 2002, 09:23:27 ; Search time 25 Seconds
(without alignments)
69.198 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10
Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 562222 seqs, 172994929 residues
Total number of hits satisfying chosen parameters: 562222

Minimum DB seq length: 0
Maximum DB seq length: 2000000000
Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : SPTREMBL19:*
1: sp-archaea:*
2: sp-bacteria:*
3: sp-fungi:*
4: sp-human:*
5: sp-invertebrate:*
6: sp-mammal:*
7: sp-mhc:*
8: sp-organelle:*
9: sp-phage:*
10: sp-plant:*
11: sp-rodent:*
12: sp-virus:*
13: sp-vertebrate:*
14: sp-unclassified:*
15: sp-rvirus:*
16: sp-bacteriap:*
17: sp-archeap:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES					Query		Match		Description	
Result No.	Score	Query	Match	Length	DB	ID				
1	55	100.0	28	4	Q9UCD1					Q9ucd1 homo sapien
2	55	100.0	30	4	Q9UCA9					Q9uca9 homo sapien
3	55	100.0	33	4	Q9UC33					Q9uc33 homo sapien
4	55	100.0	82	4	P78438					P78438 homo sapien
5	55	100.0	82	4	Q16014					Q16014 homo sapien
6	55	100.0	82	4	Q16019					Q16019 homo sapien
7	55	100.0	82	4	Q16020					Q16020 homo sapien
8	55	100.0	97	4	Q13778					Q13778 homo sapien
9	55	100.0	534	13	Q93296					Q93296 gallus gall
10	55	100.0	569	13	Q9PVL1					Q9pvl1 gallus gall
11	55	100.0	693	13	Q98SG0					Q98sg0 xenopus lae
12	55	100.0	695	6	Q95KN7					Q95kn7 macaca fasc
13	55	100.0	695	11	Q60496					Q60496 cavia sp. p
14	55	100.0	695	13	Q9DGJ8					Q9dgj8 gallus gall
15	55	100.0	747	13	Q91963					Q91963 xenopus. ap
16	55	100.0	751	13	Q9DGJ7					Q9dgj7 gallus gall

17	55	100.0	770	6	Q9TUI0					Q9tui0 sus scrofa
18	55	100.0	780	13	O73683					O73683 tetraodon f
19	52	94.5	695	13	Q9SFF9					Q9sff9 xenopus lae
20	49	89.1	612	13	Q919E7					Q919e7 brachydanio
21	49	89.1	738	13	Q90W28					Q90w28 brachydanio
22	47	85.5	79	11	O35463					O35463 cricetus
23	47	85.5	607	11	Q99K32					Q99k32 mus musculu
24	47	85.5	695	11	P97487					P97487 mus musculu
25	46	83.6	699	13	O57394					O57394 narka japon
26	46	83.6	737	13	Q93279					Q93279 fugu rubrip
27	40	72.7	19	4	Q9UCC8					Q9ucc8 homo sapien
28	39	70.9	1145	5	Q965N2					Q965n2 caenorhabdi
29	38	69.1	272	16	P96882					P96882 mycobacteri
30	38	69.1	326	2	Q9K376					Q9k376 escherichia
31	38	69.1	326	2	Q9K328					Q9k328 escherichia
32	38	69.1	326	2	Q9K2T3					Q9k2t3 escherichia
33	38	69.1	326	2	Q9K2R7					Q9k2r7 escherichia
34	38	69.1	326	2	Q9K2Q3					Q9k2q3 escherichia
35	38	69.1	326	2	Q9KH87					Q9kh87 escherichia
36	38	69.1	326	2	Q9KH85					Q9kh85 escherichia
37	38	69.1	326	2	Q9KH84					Q9kh84 escherichia
38	37	67.3	191	10	Q9SN52					Q9sn52 arabidopsis
39	37	67.3	584	5	Q9U0M8					Q9u0m8 plasmodium
40	37	67.3	1035	2	Q93E19					Q93e19 acinetobact
41	36	65.5	103	6	Q9XST6					Q9xst6 canis famil
42	36	65.5	152	11	Q9CUV7					Q9cuv7 mus musculu
43	36	65.5	190	11	Q9CPW6					Q9cpw6 mus musculu
44	36	65.5	204	11	Q9DBC2					Q9dbc2 mus musculu
45	36	65.5	210	11	Q9CQ88					Q9cq88 mus musculu
46	36	65.5	226	11	Q9CZ16					Q9cz16 mus musculu
47	36	65.5	396	4	Q9UL10					Q9ul10 homo sapien
48	36	65.5	535	3	Q01165					Q01165 magnaporthe
49	36	65.5	859	17	O26556					O26556 methanother
50	36	65.5	1668	17	O27011					O27011 methanother

ALIGNMENTS

RESULT 1

ID Q9UCD1 PRELIMINARY; PRT; 28 AA.
AC Q9UCD1;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE.
RX MEDLINE=94045685; PubMed=8229004;
RA Vigo-Pelfrey C., Lee D., Keim P., Lieberburg I., Schenk D.B.;
RT "Characterization of beta-amyloid peptide from human cerebrospinal
RT fluid.";
RL J. Neurochem. 61:1965-1968(1993).
DR HSSP; P05067; IAMB.
SQ SEQUENCE 28 AA; 3244 MW; DE7BD081160AFC81 CRC64;

Query Match 100.0%; Score 55; DB 4; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00049;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10

|||||
Db 13 HHQKLVFFAE 22

RESULT 2

Q9UCA9 PRELIMINARY; PRT; 30 AA.
ID Q9UCA9

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AC Q9UCA9;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DE 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DE 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
DE BETA-AMYLOID PROTEIN (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE
RX MEDLINE=94153015; PubMed=8109908;
RA Wisniewski T., Lalowski M., Levy E., Marques M.R., Frangione B.;
RT "The amino acid sequence of neuritic plaque amyloid from a familial
RT Alzheimer's disease patient."
RL Ann. Neurol. 35:245-246(1994).
DR HSP: P05067; IBA4.
SQ SEQUENCE 30 AA; 3391 MW; FF4167ABD081160A CRC64;

Query Match 100.0%; Score 55; DB 4; Length 30;
Best Local Similarity 100.0%; Pred. No. 0.00053;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db |||||
13 HHQKLVFFAE 22

RESULT 3
Q9UC33
ID Q9UC33 PRELIMINARY; PRT; 33 AA.
AC Q9UC33;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DE 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE
RX MEDLINE=93024877; PubMed=1406936;
RA Seubert P., Vigo-Pelfrey C., Esch F., Lee M., Dovey H., Davis D.,
RA Sinha S., Schlossmacher M., Whaley J., Swindlehurst C.;
RT "Isolation and quantification of soluble Alzheimer's beta-peptide from
RT biological fluids."
RL Nature 359:325-327(1992).
DR HSP: P05067; IBA4.
SQ SEQUENCE 33 AA; 3674 MW; BIDEFE2F4167ABD0 CRC64;

Query Match 100.0%; Score 55; DB 4; Length 33;
Best Local Similarity 100.0%; Pred. No. 0.00058;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db |||||
13 HHQKLVFFAE 22

RESULT 4
P78438
ID P78438 PRELIMINARY; PRT; 82 AA.
AC P78438;
DT 01-MAY-1997 (TrEMBLrel. 03, Created)
DT 01-MAY-1997 (TrEMBLrel. 03, Last sequence update)
DE 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
DE AMYLOID PROTEIN (BETA-AMYLOID PROTEIN) (FRAGMENT).
GN APP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
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```
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=89392030; PubMed=2675837;
RA Johnstone E.M., Chaney M.O., Moore R.E., Ward K.E., Norris F.H.,
RA Little S.P.;
RT "Alzheimer's disease amyloid peptide is encoded by two exons and shows
RT similarity to soybean trypsin inhibitor."
RL Biochem. Biophys. Res. Commun. 163:1248-1255(1989).
RN [2]
RP SEQUENCE OF 19-48 FROM N.A.
RX MEDLINE=87120329; PubMed=2949367;
RA Tanzi R.E., Gusella J.F., Watkins P.C., Bruns G.A., George-Hyslop P.,
RA Van Keuren M.L., Patterson D., Pagan S., Kurnit D.M., Neve R.L.;
RT "Amyloid beta protein gene: cDNA, mRNA distribution, and genetic
RT linkage near the Alzheimer locus."
RL Science 235:880-884(1987).
RN [3]
RP SEQUENCE OF 32-63 FROM N.A.
RX MEDLINE=93035397; PubMed=1415269;
RA Kamino K., Orr H.T., Payami H., Wijsman E.M., Alonso M.E., Pulst S.M.,
RA Anderson L., O'dahl S., Nemens E., White J.A.;
RT "Linkage and mutational analysis of familial Alzheimer disease
RT kindreds for the APP gene region."
RL Am. J. Hum. Genet. 51:998-1014(1992).
DR EMBL; M29270; AAA51768.1; -.
DR EMBL; M29269; AAA51768.1; JOINED.
DR EMBL; M15532; AAA51564.1; -.
DR EMBL; S45136; AAB23646.1; -.
DR HSP: P05067; IBA4.
FT NON_TER 1
SQ SEQUENCE 82 AA; 8994 MW; 8DA9E42B813A070E CRC64;

Query Match 100.0%; Score 55; DB 4; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.0014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db |||||
29 HHQKLVFFAE 38

RESULT 5
Q16014
ID Q16014 PRELIMINARY; PRT; 82 AA.
AC Q16014;
DT 01-NOV-1996 (TrEMBLrel. 01, Created)
DT 01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=93236601; PubMed=8476439;
RA Denman R.B., Rosenczwaig R., Miller D.L.;
RT "A system for studying the effect(s) of familial Alzheimer disease
RT mutations on the processing of the beta-amyloid peptide precursor."
RL Biochem. Biophys. Res. Commun. 192:96-103(1993).
DR EMBL; S60721; AAB26263.2; -.
DR HSP: P05067; IBA4.
FT NON_TER 1
FT NON_TER 82
SQ SEQUENCE 82 AA; 8972 MW; F534AA5B3EA9230A CRC64;

Query Match 100.0%; Score 55; DB 4; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.0014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db |||||
30 HHQKLVFFAE 39
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RESULT 6
Q16019
ID Q16019 PRELIMINARY; PRT; 82 AA.
AC Q16019;
DT 01-NOV-1996 (TREMBLrel. 01, Created)
DT 01-NOV-1996 (TREMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
GN BETA APP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=93236601; PubMed=8476439;
RA Denman R.B., Rosenzwaig R., Miller D.L.;
RT "A system for studying the effect(s) of familial Alzheimer disease
mutations on the processing of the beta-amyloid peptide precursor.";
RL Biochem. Biophys. Res. Commun. 192:96-103(1993).
DR EMBL: S61380; AAB26264.2; -
DR HSSP: P05067; 1BA4.
DR NON_TER 1
FT NON_TER 1
SQ SEQUENCE 82 AA; 8938 MW; F534AA50E579230A CRC64;

Query Match 100.0%; Score 55; DB 4; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.0014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 30 HHQKLVFFAE 39

RESULT 7
Q16020
ID Q16020 PRELIMINARY; PRT; 82 AA.
AC Q16020;
DT 01-NOV-1996 (TREMBLrel. 01, Created)
DT 01-NOV-1996 (TREMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
GN BETA APP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=93236601; PubMed=8476439;
RA Denman R.B., Rosenzwaig R., Miller D.L.;
RT "A system for studying the effect(s) of familial Alzheimer disease
mutations on the processing of the beta-amyloid peptide precursor.";
RL Biochem. Biophys. Res. Commun. 192:96-103(1993).
DR EMBL: S61383; AAB26265.2; -
DR HSSP: P05067; 1BA4.
DR NON_TER 1
FT NON_TER 1
SQ SEQUENCE 82 AA; 8882 MW; F534AA5AE5D9230A CRC64;

Query Match 100.0%; Score 55; DB 4; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.0014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 30 HHQKLVFFAE 39

RESULT 8
Q16021
ID Q16021 PRELIMINARY; PRT; 82 AA.
AC Q16021;
DT 01-NOV-1996 (TREMBLrel. 01, Created)
DT 01-NOV-1996 (TREMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
DE BETA-AMYLOID PEPTIDE (FRAGMENT).
GN BETA APP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=93236601; PubMed=8476439;
RA Denman R.B., Rosenzwaig R., Miller D.L.;
RT "A system for studying the effect(s) of familial Alzheimer disease
mutations on the processing of the beta-amyloid peptide precursor.";
RL Biochem. Biophys. Res. Commun. 192:96-103(1993).
DR EMBL: S61383; AAB26265.2; -
DR HSSP: P05067; 1BA4.
DR NON_TER 1
FT NON_TER 1
SQ SEQUENCE 82 AA; 8882 MW; F534AA5AE5D9230A CRC64;

Query Match 100.0%; Score 55; DB 4; Length 82;
Best Local Similarity 100.0%; Pred. No. 0.0014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 30 HHQKLVFFAE 39

RESULT 9
Q16022
ID Q16022 PRELIMINARY; PRT; 534 AA.
AC Q16022;
DT 01-NOV-1998 (TREMBLrel. 08, Created)
DT 01-NOV-1998 (TREMBLrel. 08, Last sequence update)
DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
DE AMYLOID PROTEIN (FRAGMENT).
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianinae;
OC Gallus.
OX NCBI_TaxID=9031;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=98337885; PubMed=9671674;
RA Barnes N.Y., Li L., Yoshikawa K., Schwartz L.M., Oppenheim R.W.,
RA Milligan C.E.;
RT "Increased production of amyloid precursor protein provides a
substrate for caspase-3 in dying motoneurons.";
RL J. Neurosci. 18:5869-5880(1998).
DR EMBL: AF042098; AAC25052.1; -
DR HSSP: P05067; 1BA4.
DR InterPro: IPR001868; A4_APP.
DR PRINTS: PR00203; AMYLOIDA4.
DR PROSITE: PS00319; A4_EXTRA; 1.
DR PROSITE: PS00320; A4_INTRA; 1.
FT NON_TER 1
SQ SEQUENCE 534 AA; 60597 MW; FB53ECC2E66D4C92 CRC64;

Query Match 100.0%; Score 55; DB 13; Length 534;
Best Local Similarity 100.0%; Pred. No. 0.0095;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 448 HHQKLVFFAE 457

RESULT 9
Q13778
ID Q13778 PRELIMINARY; PRT; 97 AA.
AC Q13778;
DT 01-NOV-1996 (TREMBLrel. 01, Created)
DT 01-NOV-1996 (TREMBLrel. 01, Last sequence update)
DT 01-JUN-2001 (TREMBLrel. 17, Last annotation update)
DE AMYLOID PROTEIN (AD-AP) (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=87120328; PubMed=3810169;
RA Goldgaber D., Lerman M.I., McBride O.W., Saffioti U., Gajdusek D.C.;
RT "Characterization and chromosomal localization of a cDNA encoding
brain amyloid Cf Alzheimer's disease.";
RL Science 235:877-880(1987).
DR EMBL: M15533; AAA35540.1; -
DR HSSP: P05067; 1BA4.
DR InterPro: IPR001868; A4_APP.
DR PRINTS: PR00203; AMYLOIDA4.
FT NON_TER 1
SQ SEQUENCE 97 AA; 10884 MW; E528CDB448DE474E CRC64;

Query Match 100.0%; Score 55; DB 4; Length 97;
Best Local Similarity 100.0%; Pred. No. 0.0017;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 11 HHQKLVFFAE 20

RESULT 9
Q13779
ID Q13779 PRELIMINARY; PRT; 534 AA.
AC Q13779;
DT 01-NOV-1998 (TREMBLrel. 08, Created)
DT 01-NOV-1998 (TREMBLrel. 08, Last sequence update)
DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
DE AMYLOID PROTEIN (FRAGMENT).
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianinae;
OC Gallus.
OX NCBI_TaxID=9031;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=98337885; PubMed=9671674;
RA Barnes N.Y., Li L., Yoshikawa K., Schwartz L.M., Oppenheim R.W.,
RA Milligan C.E.;
RT "Increased production of amyloid precursor protein provides a
substrate for caspase-3 in dying motoneurons.";
RL J. Neurosci. 18:5869-5880(1998).
DR EMBL: AF042098; AAC25052.1; -
DR HSSP: P05067; 1BA4.
DR InterPro: IPR001868; A4_APP.
DR PRINTS: PR00203; AMYLOIDA4.
DR PROSITE: PS00319; A4_EXTRA; 1.
DR PROSITE: PS00320; A4_INTRA; 1.
FT NON_TER 1
SQ SEQUENCE 534 AA; 60597 MW; FB53ECC2E66D4C92 CRC64;

Query Match 100.0%; Score 55; DB 13; Length 534;
Best Local Similarity 100.0%; Pred. No. 0.0095;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 448 HHQKLVFFAE 457
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RESULT 10
Q9PVL1 ID Q9PVL1 PRELIMINARY; PRT; 569 AA.
AC Q9PVL1;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE AMYLOID PROTEIN (FRAGMENT).
GN APP.
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianinae;
OC Gallus.
OC NCBI_TaxID=9031;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=BRAIN;
RA Coulson E.J., Paliga K., Beyreuther K., Masters C.L.;
RT "What the evolution of the amyloid protein precursor supergene family
RT tells us about its function.";
RL Neurochem. Int. 0:0-0(2000).
DR EMBL; AF030341; AAF12698.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
FT NON_TER 1
SQ SEQUENCE 569 AA; 64753 MW; 0AB8B851863A19D CRC64;

Query Match
Best Local Similarity 100.0%; Score 55; DB 13; Length 569;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 484 HHQKLVFFAE 493
|||||

RESULT 11
Q98SGO ID Q98SGO PRELIMINARY; PRT; 693 AA.
AC Q98SGO;
DT 01-JUN-2001 (TREMBlrel. 17, Created)
DT 01-JUN-2001 (TREMBlrel. 17, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE BETA-AMYLOID PRECURSOR PROTEIN A.
GN APP.
OS Xenopus laevis (African clawed frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Mesobatrachia; Pipidae;
OC Xenopodinae; Xenopus.
OC NCBI_TaxID=8355;
RN [1]
RP SEQUENCE FROM N.A.
RA Van den Hurk W.H.;
RL Thesis (2001), Department of Biological Sciences,
RL University of Nijmegen, Nijmegen, Netherlands.
DR EMBL; AJ298150; CAC37193.1; -.
DR HSSP; P05067; 1H23.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
KW Signal.
FT SIGNAL 1 18 POTENTIAL.
SQ SEQUENCE 693 AA; 78568 MW; CAF1DF55C1AB653 CRC64;

Query Match
Best Local Similarity 100.0%; Score 55; DB 13; Length 693;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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Best Local Similarity 100.0%; Pred. No. 0.012;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 607 HHQKLVFFAE 616
|||||

RESULT 12
Q95KN7 ID Q95KN7 PRELIMINARY; PRT; 695 AA.
AC Q95KN7;
DT 01-DEC-2001 (TREMBlrel. 19, Created)
DT 01-DEC-2001 (TREMBlrel. 19, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE AMYLOID B-PROTEIN PRECURSOR.
OS Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Cercopithecoidea;
OC Cercopithecinae; Macaca.
OC NCBI_TaxID=9541;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=CEREBELLUM;
RX MEDLINE=91273117; PubMed=1905108;
RA Podlisky M.B., Toland D.R., Selkoe D.J.;
RT "Homology of the amyloid beta protein precursor in monkey and human
RT supports a primate model for beta amyloidosis in Alzheimer's
RT disease.";
RL Am. J. Pathol. 138:1423-1435(1991).
DR EMBL; M58727; AAA36829.1; -. POTENTIAL.
FT SIGNAL 1 17
FT CHAIN 597 636 POTENTIAL.
SQ SEQUENCE 695 AA; 78663 MW; 4F6EA0139F969D56 CRC64;

Query Match
Best Local Similarity 100.0%; Score 55; DB 6; Length 695;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 509 HHQKLVFFAE 618
|||||

RESULT 13
Q60496 ID Q60496 PRELIMINARY; PRT; 695 AA.
AC Q60496;
DT 01-NOV-1996 (TREMBlrel. 01, Created)
DT 01-NOV-1996 (TREMBlrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE PUTATIVE AMYLOID PRECURSOR PROTEIN.
OS Cavia sp.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Hystricognathi; Caviidae; Cavia.
OC NCBI_TaxID=10143;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=BRAIN;
RX MEDLINE=97236426; PubMed=9116031;
RA Beck M., Mueller D., Bigl V.;
RT "Amyloid precursor protein in Guinea pigs - complete cDNA sequence and
RT alternative splicing.";
RL Biochim. Biophys. Acta 1351:17-21(1997).
DR EMBL; X97631; CAA66230.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
SQ SEQUENCE 695 AA; 78701 MW; 5196A0C4017F16AB CRC64;

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Query Match      100.0%; Score 55; DB 11; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.012;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
   |
Db 609 HHQKLVFFAE 618
   |

RESULT 14
Q9DGS8          PRELIMINARY;      PRT;      695 AA.
AC Q9DGS8;
DT 01-MAR-2001 (Tremblrel. 16, Created)
DT 01-MAR-2001 (Tremblrel. 16, Last sequence update)
DT 01-DEC-2001 (Tremblrel. 19, Last annotation update)
DE BETA-AMYLOID PRECURSOR PROTEIN 695 ISOFORM.
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
OC Gallus.
OX NCBI_TaxID=9031;
RN [1]
RP SEQUENCE FROM N.A.
RA Sarasa M., Rodolosse A., Sorribas V.;
RT "Cloning of full-length chicken beta-amyloid precursor protein
RT isoforms.";
RL Submitted (JUL-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF289218; AAC00593.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
SQ SEQUENCE 695 AA; 78565 MW; F201ED02AEC86D95 CRC64;

Query Match      100.0%; Score 55; DB 13; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.012;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
   |
Db 609 HHQKLVFFAE 618
   |

RESULT 15
Q91963          PRELIMINARY;      PRT;      747 AA.
AC Q91963;
DT 01-NOV-1996 (Tremblrel. 01, Created)
DT 01-NOV-1996 (Tremblrel. 01, Last sequence update)
DT 01-DEC-2001 (Tremblrel. 19, Last annotation update)
DE APP747.
GN APP747.
OS Xenopus.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Mesobatrachia; Pipidea; Pipidae;
OC Xenopodinae.
OX NCBI_TaxID=8353;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE-93129227; PubMed-1282805;
RA Okado H., Okamoto H.;
RT "A Xenopus homologue of the human beta-amyloid precursor protein:
RT developmental regulation of its gene expression.";
RL Biochem. Biophys. Res. Commun. 189:1561-1568(1992).
DR EMBL; S52417; AAB24853.1; -.
DR HSSP; P05067; 1HZ3.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
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DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00006; A4_EXTRA; 1.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS50279; BPTI_KUNITZ_2; 1.
KW Serine protease inhibitor.
SQ SEQUENCE 747 AA; 84893 MW; A75E81885681D948 CRC64;

Query Match      100.0%; Score 55; DB 13; Length 747;
Best Local Similarity 100.0%; Pred. No. 0.013;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
   |
Db 661 HHQKLVFFAE 670
   |

RESULT 16
Q9DGS7          PRELIMINARY;      PRT;      751 AA.
AC Q9DGS7;
DT 01-MAR-2001 (Tremblrel. 16, Created)
DT 01-MAR-2001 (Tremblrel. 16, Last sequence update)
DT 01-DEC-2001 (Tremblrel. 19, Last annotation update)
DE BETA-AMYLOID PRECURSOR PROTEIN 751 ISOFORM.
OS Gallus gallus (Chicken).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
OC Gallus.
OX NCBI_TaxID=9031;
RN [1]
RP SEQUENCE FROM N.A.
RA Sarasa M., Rodolosse A., Sorribas V.;
RT "Cloning of full-length chicken beta-amyloid precursor protein
RT isoforms.";
RL Submitted (JUL-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF289219; AAC00594.1; -.
DR HSSP; P05067; 1BA4.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00006; A4_EXTRA; 1.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS50279; BPTI_KUNITZ_2; 1.
KW Serine protease inhibitor.
SQ SEQUENCE 751 AA; 84705 MW; E78E9413A8033D84 CRC64;

Query Match      100.0%; Score 55; DB 13; Length 751;
Best Local Similarity 100.0%; Pred. No. 0.013;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
   |
Db 665 HHQKLVFFAE 674
   |

RESULT 17
Q9TU10          PRELIMINARY;      PRT;      770 AA.
AC Q9TU10;
DT 01-MAY-2000 (Tremblrel. 13, Created)
DT 01-MAY-2000 (Tremblrel. 13, Last sequence update)
DT 01-DEC-2001 (Tremblrel. 19, Last annotation update)
DE AMYLOID PRECURSOR PROTEIN.
OS Sus scrofa (pig).
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OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
OX NCBI_TaxID=9823;
RN [1]
RP SEQUENCE FROM N.A.
RA Kimura A., Takahashi T.;
RT "Amyloid Precursor Protein 770."
RL Submitted (SEP-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AB032550; BAA84580.1; -.
DR HSSP; P05067; 1AAP.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRASE.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS0279; BPTI_KUNITZ_2; 1.
DR Serine protease inhibitor.
KW Glycoprotein; Amyloid; Neurone; Transmembrane; Signal;
FT SIGNAL 1 18
FT CHAIN 19 780
FT FT
FT CHAIN 682 724
FT DOMAIN 19 711
FT TRANSMEM 712 732
FT DOMAIN 733 780
FT DOMAIN 323 382
FT SITE 769 772
FT DISULFID 327 378
FT DISULFID 336 361
FT CARBOHYD 560 560
SQ SEQUENCE 770 AA; 86961 MW; 5F7A1DCB2BCC583E CRC64;

Query Match 100.0%; Score 55; DB 6; Length 770;
Best Local Similarity 100.0%; Pred. No. 0.014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 684 HHQKLVFFAE 693
|||||

RESULT 18
O73683
ID O73683 PRELIMINARY; PRT; 780 AA.
AC O73683;
DT 01-AUG-1998 (TrEMBLrel. 07, Created)
DT 01-AUG-1998 (TrEMBLrel. 07, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE ALZHEIMER'S DISEASE AMYLOID A4 PROTEIN HOMOLOG PRECURSOR [CONTAINS:
DE BETA-AMYLOID PROTEIN (BETA-APP) (A-BETA)].
GN APP.
OS Tetraodon fluviatilis (Puffer fish).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC Acanthomorpha; Acanthopterygii; Percomorpha; Tetraodontiformes;
OC Tetraodontidae; Tetraodon.
OX NCBI_TaxID=47145;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE-98252138; PubMed-9599080;
RA Villard L., Tassone F., Crnogorac-Jurcevic T., Clancy K., Gardiner K.;
RT "Analysis of pufferfish homologues of the AT-rich human APP gene.";
RL Gene 210:17-24(1998).
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
CC INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN
CC G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: TYPE I MEMBRANE PROTEIN.
CC -!- DOMAIN: THE CLATHRIN-BINDING SITE IS ESSENTIAL FOR ITS ASSOCIATION
CC WITH X11-ALPHA, -BETA, AND -GAMMA. THE SEQUENCE SPECIFIC
CC RECOGNITION EXTENDS TO PEPTIDE RESIDUES THAT ARE C-TERMINAL TO THE
CC NPYX MOTIF. THIS INTERACTION APPEARS TO BE INDEPENDENT OF
CC PHOSPHORYLATION (BY SIMILARITY).
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -!- SIMILARITY: CONTAINS 1 PROTEASE INHIBITOR DOMAIN BELONGING TO THE
CC BPTI/KUNITZ FAMILY OF INHIBITORS.
DR EMBL; AF018165; AAC41275.1; -.
DR HSSP; P05067; 1HZ3.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
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DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00006; A4_EXTRA; 1.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; FALSE_NEG.
DR PROSITE; PS0279; BPTI_KUNITZ_2; 1.
KW Glycoprotein; Amyloid; Neurone; Transmembrane; Signal;
KW serine protease inhibitor.
FT SIGNAL 1 18
FT CHAIN 19 780
FT FT
FT CHAIN 682 724
FT DOMAIN 19 711
FT TRANSMEM 712 732
FT DOMAIN 733 780
FT DOMAIN 323 382
FT SITE 769 772
FT DISULFID 327 378
FT DISULFID 336 361
FT CARBOHYD 560 560
SQ SEQUENCE 780 AA; 88238 MW; 60071BE94520191D CRC64;

Query Match 100.0%; Score 55; DB 13; Length 780;
Best Local Similarity 100.0%; Pred. No. 0.014;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 694 HHQKLVFFAE 703
|||||

RESULT 19
Q98SF9
ID Q98SF9 PRELIMINARY; PRT; 695 AA.
AC Q98SF9;
DT 01-JUN-2001 (TrEMBLrel. 17, Created)
DT 01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE BETA-AMYLOID PRECURSOR PROTEIN B.
GN APP.
OS Xenopus laevis (African clawed frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Mesobatrachia; Pipoidae; Pipidae;
OC Xenopodinae; Xenopus.
OX NCBI_TaxID=8355;
RN [1]
RP SEQUENCE FROM N.A.
RA Van den Hurk W.H.;
RL Thesis (2001), Department of Biological Sciences,
RL University of Nijmegen, Nijmegen, Netherlands.
DR EMBL; AJ298151; CAC37194.1; -.
DR HSSP; P05067; 1HZ3.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
KW Signal.
FT SIGNAL 1 18
FT CHAIN 19 780
FT FT
FT CHAIN 682 724
FT DOMAIN 19 711
FT TRANSMEM 712 732
FT DOMAIN 733 780
FT DOMAIN 323 382
FT SITE 769 772
FT DISULFID 327 378
FT DISULFID 336 361
FT CARBOHYD 560 560
SQ SEQUENCE 780 AA; 88238 MW; 60071BE94520191D CRC64;

Query Match 94.5%; Score 52; DB 13; Length 695;
Best Local Similarity 90.0%; Pred. No. 0.049;
Matches 9; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
Db 609 HHQKLVFFAD 618
|||||
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RESULT 20
Q919E7 PRELIMINARY; PRT; 612 AA.
AC Q919E7
DT 01-OCT-2000 (TReMBLrel. 15, Created)
DT 01-OCT-2000 (TReMBLrel. 15, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE AMYLOID PROTEIN (FRAGMENT)
OS Brachydanio rerio (zebrafish) (Zebra danio).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Ostariophysi;
OC Cypriniformes; Cyprinidae; Danio.
OX NCBI_TaxID=7955;
RN [1]
RP SEQUENCE FROM N.A.
RA Slavov D.B., Gardiner K.;
RT "An App cDNA from zebrafish (Danio rerio).";
RL Submitted (APR-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF257742; AAF71748.1; -
DR HSSP; P05067; 1H23.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
FT NON_TER 1
SQ SEQUENCE 612 AA; 69710 MW; 59A9ACBDF9C59EFF CRC64;

Query Match 89.1%; Score 49; DB 13; Length 612;
Best Local Similarity 90.0%; Pred. No. 0.17;
Matches 9; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 526 YHQLVFFAE 535

RESULT 21
Q90W28 PRELIMINARY; PRT; 738 AA.
AC Q90W28;
DT 01-DEC-2001 (TReMBLrel. 19, Created)
DT 01-DEC-2001 (TReMBLrel. 19, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE AMYLOID PRECURSOR PROTEIN.
GN APP.
OS Brachydanio rerio (zebrafish) (Zebra danio).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Ostariophysi;
OC Cypriniformes; Cyprinidae; Danio.
OX NCBI_TaxID=7955;
RN [1]
RP SEQUENCE FROM N.A.
RA Groth C., Lardelli M.;
RT "Expression analysis of zebrafish app.";
RL Submitted (JUN-2001) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF389401; AAK64495.1; -
SQ SEQUENCE 738 AA; 83577 MW; AF480F6D308FD298 CRC64;

Query Match 89.1%; Score 49; DB 13; Length 738;
Best Local Similarity 90.0%; Pred. No. 0.2;
Matches 9; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
DB 652 YHQLVFFAE 661

RESULT 22
Q35463 PRELIMINARY; PRT; 79 AA.
ID O35463
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AC O35463;
DT 01-JAN-1998 (TReMBLrel. 05, Created)
DT 01-JAN-1998 (TReMBLrel. 05, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE ALZHEIMER'S AMYLOID BETA PROTEIN (FRAGMENT).
GN BETA APP.
OS Cricetus griseus (Chinese hamster).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Cricetinae;
OC Cricetulus.
OX NCBI_TaxID=10029;
RN [1]
RP SEQUENCE FROM N.A.
RA Sambamurti K., Pinnix I., Gandhi S.;
RL Submitted (OCT-1997) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF030413; AAB66608.1; -
DR HSSP; P05067; 1BA4.
FT NON_TER 1
FT NON_TER 79
SQ SEQUENCE 79 AA; 8538 MW; 37F2C6C3BFF3F597 CRC64;

Query Match 85.5%; Score 47; DB 11; Length 79;
Best Local Similarity 100.0%; Pred. No. 0.053;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFAE 10
DB 34 HOKLVFFAE 42

RESULT 23
Q99K32 PRELIMINARY; PRT; 607 AA.
ID Q99K32;
AC Q99K32;
DT 01-JUN-2001 (TReMBLrel. 17, Created)
DT 01-JUN-2001 (TReMBLrel. 17, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE HYPOTHETICAL 68.4 KDA PROTEIN (FRAGMENT).
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RA TISSUE-MAMMARY TUMOR. WAP-TGF ALPHA MODEL. 7 MONTHS OLD, GROSS
RC TISSUE.;
RA Strausberg R.;
RL Submitted (MAR-2001) to the EMBL/GenBank/DBJ databases.
DR EMBL; BC005490; AAH05490.1; -
DR HSSP; P05067; 1AAP.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS02079; BPTI_KUNITZ_2; 1.
KW Hypothetical protein; Serine protease inhibitor.
FT NON_TER 1
SQ SEQUENCE 607 AA; 68391 MW; BF802214CBA7D172 CRC64;

Query Match 85.5%; Score 47; DB 11; Length 607;
Best Local Similarity 100.0%; Pred. No. 0.42;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HOKLVFFAE 10
DB 522 HOKLVFFAE 530
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RESULT 24
P97487 PRELIMINARY; PRT; 695 AA.
AC P97487;
DT 01-MAY-1997 (TrEMBLrel. 03, Created)
DT 01-MAY-1997 (TrEMBLrel. 03, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE HIPPOCAMPAL AMYLOID PROTEIN.
GN APP.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RN SEQUENCE FROM N.A.
RC STRAIN=SAMP8; TISSUE=HIPPOCAMPUS;
RA Flood J.F., Kumar V.B., Sasser T., Word I., Morley J.E.;
RL Submitted (JAN-1997) to the EMBL/GenBank/DBJ databases.
RN [2]
RN SEQUENCE OF 581-662 FROM N.A.
RC STRAIN=129SV;
RA Wragg M.A., Busfield F., Duff K., Korenblat K., Capocchi M.,
RA Loring J.F., Coate A.M.;
RL Submitted (DEC-1996) to the EMBL/GenBank/DBJ databases.
DE EMBL; U84012; ABA41502.1; -.
DR EMBL; U82624; ABA40919.1; -.
DR HSSP; P05067; IMWP.
DR MGD; MGI:88059; App.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
SQ SEQUENCE 695 AA; 78414 MW; 9A5FBE2ED261236E CRC64;

Query Match 85.5%; Score 47; DB 11; Length 695;
Best Local Similarity 100.0%; Pred. No. 0.48;
Matches 9; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQKLVEFAE 10
| | | | | | | |
DB 610 HQKLVEFAE 618

RESULT 25
O57394 PRELIMINARY; PRT; 699 AA.
AC O57394;
DT 01-JUN-1998 (TrEMBLrel. 06, Created)
DT 01-JUN-1998 (TrEMBLrel. 06, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE EL AMYLOID PRECURSOR PROTEIN 699.
GN EL APP699.
OS Narke japonica (Electric ray).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Chondrichthyes;
OC Elasmobranchii; Squala; Hypnosqualea; Pristiogaster; Batoidae;
OC Torpediniformes; Narcineidae; Naridae; Narke.
OX NCBI_TaxID=62965;
RN [1]
RN SEQUENCE FROM N.A.
RC TISSUE=ELECTRIC LOBE;
RX MEDLINE=98129705; PubMed=9461486;
RA Iijima K., Lee D.-S., Okutsu J., Tomita S., Hirashima N., Kirino Y.,
RA Suzuki T.;
RT "cDNA isolation of Alzheimer's amyloid precursor protein from
RT cholinergic nerve terminals of the electric organ of the electric
RT ray.";
RL Biochem. J. 330:29-33(1998).
DR EMBL; AB005544; BAA24230.1; -.
DR HSSP; P05067; 1H23.
DR InterPro; IPR001868; A4_APP.
DR Pfam; PF02177; A4_EXTRA; 1.
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DR PRINTS; PR00203; AMYLOIDA4.
DR SMART; SM00006; A4_EXTRA; 1.
DR PROSITE; PS00319; A4_EXTRA; 1.
DR PROSITE; PS00320; A4_INTRA; 1.
SQ SEQUENCE 699 AA; 78879 MW; 952915C309D50E5C CRC64;

Query Match 83.6%; Score 46; DB 13; Length 699;
Best Local Similarity 100.0%; Pred. No. 0.76;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFF 8
| | | | | | | |
DB 613 HHOKLVFF 620

RESULT 26
O93279 PRELIMINARY; PRT; 737 AA.
AC O93279;
DT 01-NOV-1998 (TrEMBLrel. 08, Created)
DT 01-NOV-1998 (TrEMBLrel. 08, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE ALZHEIMER'S DISEASE AMYLOID A4 PROTEIN HOMOLOG PRECURSOR [CONTAINS:
DE BETA-AMYLOID PROTEIN (BETA-APP) (A-BETA)].
GN APP.
OS Fugu rubripes (Japanese pufferfish) (Takifugu rubripes).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC Acanthomorpha; Acanthopterygii; Percomorpha; Tetraodontiformes;
OC Tetraodontidae; Takifugu.
OX NCBI_TaxID=31033;
RN [1]
RN SEQUENCE FROM N.A.
RX MEDLINE=98252138; PubMed=9599080;
RA Villard L., Tassone F., Crnogorac-Jurcovic T., Clancy K., Gardiner K.;
RT "Analysis of pufferfish homologues of the A4-rich human APP gene.";
CC -!- FUNCTION: FUNCTIONAL NEURONAL RECEPTOR WHICH COUPLES TO
CC INTRACELLULAR SIGNALING PATHWAY THROUGH THE GTP-BINDING PROTEIN
CC G(O) (BY SIMILARITY).
CC -!- SUBCELLULAR LOCATION: TYPE I MEMBRANE PROTEIN.
CC -!- DOMAIN: THE CLATHRIN-BINDING SITE IS ESSENTIAL FOR ITS ASSOCIATION
CC WITH X11-ALPHA, -BETA, AND -GAMMA. THE SEQUENCE SPECIFIC
CC RECOGNITION EXTENDS TO PEPTIDE RESIDUES THAT ARE C-TERMINAL TO THE
CC NPXY MOTIF. THIS INTERACTION APPEARS TO BE INDEPENDENT OF
CC PHOSPHORYLATION (BY SIMILARITY).
CC -!- SIMILARITY: BELONGS TO THE APP FAMILY.
CC -!- SIMILARITY: CONTAINS 1 PROTEASE INHIBITOR DOMAIN BELONGING TO THE
CC BPTI/KUNITZ FAMILY OF INHIBITORS.
DR EMBL; AF090120; AAD13392.1; -.
DR HSSP; P05067; 1H23.
DR InterPro; IPR001868; A4_APP.
DR InterPro; IPR002223; Kunitz_BPTI.
DR Pfam; PF02177; A4_EXTRA; 1.
DR Pfam; PF00014; Kunitz_BPTI; 1.
DR PRINTS; PR00203; AMYLOIDA4.
DR PRINTS; PR00759; BASICPTASE.
DR SMART; SM00006; A4_EXTRA; 1.
DR SMART; SM00131; KU; 1.
DR PROSITE; PS00319; A4_EXTRA; FALSE_NEG.
DR PROSITE; PS00320; A4_INTRA; 1.
DR PROSITE; PS00280; BPTI_KUNITZ_1; 1.
DR PROSITE; PS00279; BPTI_KUNITZ_2; 1.
KW Glycoprotein; Amyloid; Neurone; Transmembrane; Signal;
KW Serine protease inhibitor.
FT SIGNAL 1 18 POTENTIAL.
FT CHAIN 19 737 ALZHEIMER'S DISEASE AMYLOID A4 PROTEIN
FT HOMOLOG.
FT CHAIN 639 681 BETA-AMYLOID PROTEIN (POTENTIAL).
FT DOMAIN 19 668 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 669 689 POTENTIAL.
FT DOMAIN 690 737 CYTOPLASMIC (POTENTIAL).
FT DOMAIN 286 344 BPTI/KUNITZ INHIBITOR.
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FT SITE 726 729 CLATHRIN-BINDING (BY SIMILARITY).
FT ACT_SITE 300 301 REACTIVE BOND.
FT DISULFID 290 340 BY SIMILARITY.
FT DISULFID 299 323 BY SIMILARITY.
FT DISULFID 315 336 BY SIMILARITY.
FT CARBOHYD 522 522 N-LINKED (GLCNAC...) (POTENTIAL).
SQ SEQUENCE 737 AA; 82856 MW; 6PAD01E2E3B2B7E2 CRC64;

Query Match 83.6%; Score 46; DB 13; Length 737;
Best Local Similarity 80.0%; Pred. No. 0.8;
Matches 8; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
Db 651 YHQLVFFAD 660
:|||||||

RESULT 27
Q9UCC8 PRELIMINARY; PRT; 19 AA.
AC Q9UCC8;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DE BETA-AMYLOID-(1-42) (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE.
RX MEDLINE=94068497; PubMed=8248178;
RA Rober A.E., Lovenson J.D., Clarke S., Woods A.S., Cotter R.J.,
RA Gowling E., Ball M.J.;
RT "beta-Amyloid-(1-42) is a major component of cerebrovascular amyloid
RT deposits: implications for the pathology of Alzheimer disease.";
RL Proc. Natl. Acad. Sci. U.S.A. 90:10836-10840(1993).
DR HSSP: P05067; 1AMB.
SQ SEQUENCE 19 AA; 2315 MW; 05B02B3F6DDECE3E CRC64;

Query Match 72.7%; Score 40; DB 4; Length 19;
Best Local Similarity 100.0%; Pred. No. 0.31;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVF 7
Db 13 HHQKLVF 19
|||||||

RESULT 28
Q965N2 PRELIMINARY; PRT; 1145 AA.
AC Q965N2;
DT 01-DEC-2001 (TREMBlrel. 19, Created)
DT 01-DEC-2001 (TREMBlrel. 19, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE HYPOTHETICAL PROTEIN BE0003N10.3.
GN BE0003N10.3.
OS Caenorhabditis elegans.
OC Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida; Rhabditoidea;
OC Rhabditidae; Peloderinae; Caenorhabditis.
OX NCBI_TaxID=6239;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN-BRISTOL N2;
RX MEDLINE=99069613; PubMed=9851916;
RA None;
RT "Genome sequence of the nematode C. elegans: a platform for
RT investigating biology. The C. elegans Sequencing Consortium.";
RL Science 282:2012-2018(1998).
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN-BRISTOL N2;

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RA Waterston R.;
RT "Direct Submission.";
RL Submitted (AUG-2001) to the EMBL/GenBank/DBJ databases.
DR EMBL; AC092690; AAK73857.1; -.
SQ SEQUENCE 1145 AA; 128815 MW; 67EC2437F4F4A377 CRC64;

Query Match 70.9%; Score 39; DB 5; Length 1145;
Best Local Similarity 60.0%; Pred. No. 30;
Matches 6; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
Db 512 HHEKLFLHE 521
|||||

RESULT 29
P96882 PRELIMINARY; PRT; 272 AA.
AC P96882;
DT 01-MAY-1997 (TREMBlrel. 03, Created)
DT 01-MAY-1997 (TREMBlrel. 03, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE HYPOTHETICAL 30.1 KDA PROTEIN.
GN RV3277 OR MTCY71.17.
OS Mycobacterium tuberculosis.
OC Bacteria; Firmicutes; Actinobacteria; Actinobacteridae;
OC Actinomycetales; Corynebacterineae; Mycobacteriaceae; Mycobacterium.
OX NCBI_TaxID=1773;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=H37RV;
RX MEDLINE=98295987; PubMed=9634230;
RA Cole S.T., Brosch R., Parkhill J., Garnier T., Churcher C., Harris D.,
RA Gordon S.V., Eigmeier K., Gas S., Barry C.E. III, Tekala F.,
RA Badcock K., Basham D., Brown D., Chillingworth T., Connor R.,
RA Davies R., Devlin K., Feltwell T., Gentles S., Hamlin N., Holroyd S.,
RA Hornsby T., Jagels K., Krogh A., McLean J., Moule S., Murphy L.,
RA Oliver S., Osborne J., Quail M.A., Rajandream M.A., Rogers J.,
RA Rutter S., Seeger K., Skelton S., Squares S., Squares R.,
RA Sulston J.E., Taylor K., Whitehead S., Barrell B.G.;
RT "Deciphering the biology of Mycobacterium tuberculosis from the
RT complete genome sequence.";
RL Nature 393:537-544(1998).
DR EMBL; Z92771; CAB07080.1; -.
DR Tuberculist; RV3277; -.
KW Hypothetical protein; Complete proteome.
SQ SEQUENCE 272 AA; 30078 MW; F07597B96A0AB081 CRC64;

Query Match 69.1%; Score 38; DB 16; Length 272;
Best Local Similarity 66.7%; Pred. No. 11;
Matches 6; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

OY 1 HHQKLVFFA 9
Db 137 HHEALLFFA 145
|||||

RESULT 30
Q9K376 PRELIMINARY; PRT; 326 AA.
AC Q9K376;
DT 01-OCT-2000 (TREMBlrel. 15, Created)
DT 01-OCT-2000 (TREMBlrel. 15, Last sequence update)
DT 01-JUN-2001 (TREMBlrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=CL-3, AND DEC8B;

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RX MEDLINE=20351039; PubMed=10894541;
RT Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RL "Parallel evolution of virulence in pathogenic Escherichia coli.";
DR EMBL; AF267594; AAF97134.1; -.
DR EMBL; AF267594; AAF97127.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36326 MW; 326C60E6F59A625C CRC64;

Query Match 69.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
Db 289 HHQKLLSKFFAQ 300

RESULT 31
ID Q9K328 PRELIMINARY; PRT; 326 AA.
AC Q9K328;
DT 01-OCT-2000 (TrEMBLrel. 15, Created)
DT 01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TrEMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=921-B4, CL-37, B170, AND G5506;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RL "Parallel evolution of virulence in pathogenic Escherichia coli.";
DR EMBL; AF267597; AAF97137.1; -.
DR EMBL; AF267588; AAF97128.1; -.
DR EMBL; AF267592; AAF97132.1; -.
DR EMBL; AF267596; AAF97136.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36333 MW; 51A210B6F59A6248 CRC64;

Query Match 69.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
Db 289 HHQKLLFKFFAQ 300

RESULT 32
Q9K2T3 PRELIMINARY; PRT; 326 AA.
AC Q9K2T3;
DT 01-OCT-2000 (TrEMBLrel. 15, Created)
DT 01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TrEMBLrel. 17, Last annotation update)

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DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=DEC12A, AND DEC11A;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RL "Parallel evolution of virulence in pathogenic Escherichia coli.";
DR EMBL; AF267591; AAF97131.1; -.
DR EMBL; AF267591; AAF97130.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36347 MW; A4A740B3F0CF624E CRC64;

Query Match 69.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
Db 289 HHQKLLFKFFAQ 300

RESULT 33
Q9K2R7 PRELIMINARY; PRT; 326 AA.
AC Q9K2R7;
DT 01-OCT-2000 (TrEMBLrel. 15, Created)
DT 01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TrEMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=DEC2A, E2348/69, AND DEC1A;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RL "Parallel evolution of virulence in pathogenic Escherichia coli.";
DR EMBL; AF267581; AAF97121.1; -.
DR EMBL; AF267579; AAF97119.1; -.
DR EMBL; AF267580; AAF97120.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36400 MW; B1B240B3F0CF624E CRC64;

Query Match 69.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHQKLV--FFAE 10
Db 289 HHQKLLFKFFAQ 300

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RESULT 34
Q9K203
ID Q9K203 PRELIMINARY; PRT; 326 AA.
AC Q9K203;
DT 01-OCT-2000 (TReMBLrel. 15, Created)
DT 01-OCT-2000 (TReMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TReMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=DEC3D, 93-111, OK-1, DEC3F, AND 5905;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RT "Parallel evolution of virulence in pathogenic Escherichia coli.";
RL Nature 406:64-67(2000).
DR EMBL; AF267586; AAF97126.1; -.
DR EMBL; AF267582; AAF97122.1; -.
DR EMBL; AF267583; AAF97123.1; -.
DR EMBL; AF267584; AAF97124.1; -.
DR EMBL; AF267585; AAF97125.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36259 MW; BC10FECA2EFC1F7A CRC64;

Query Match 59.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
Db 289 HHOKLLSKFFAQ 300
|||||: |||:

RESULT 35
Q9KH87
ID Q9KH87 PRELIMINARY; PRT; 326 AA.
AC Q9KH87;
DT 01-OCT-2000 (TReMBLrel. 15, Created)
DT 01-OCT-2000 (TReMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TReMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=DEC3F;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RT "Parallel evolution of virulence in pathogenic Escherichia coli.";
RL Nature 406:64-67(2000).
DR EMBL; AF267589; AAF97129.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36319 MW; 92A210FE5690515A CRC64;

Query Match 59.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
Db 289 HHOKLLSKFFAQ 300
|||||: |||:

RESULT 36
Q9KH85
ID Q9KH85 PRELIMINARY; PRT; 326 AA.
AC Q9KH85;
DT 01-OCT-2000 (TReMBLrel. 15, Created)
DT 01-OCT-2000 (TReMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TReMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=B2F1;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RT "Parallel evolution of virulence in pathogenic Escherichia coli.";
RL Nature 406:64-67(2000).
DR EMBL; AF267595; AAF97135.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
DR PROSITE; PS00755; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36312 MW; A905FECA2EFC1F7A CRC64;

Query Match 59.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 2; Gaps 1;

QY 1 HHOKLV--FFAE 10
Db 289 HHOKLLSKFFAQ 300
|||||: |||:

RESULT 37
Q9KH84
ID Q9KH84 PRELIMINARY; PRT; 326 AA.
AC Q9KH84;
DT 01-OCT-2000 (TReMBLrel. 15, Created)
DT 01-OCT-2000 (TReMBLrel. 15, Last sequence update)
DT 01-JUN-2001 (TReMBLrel. 17, Last annotation update)
DE PHOSPHOGLUCOSE ISOMERASE (FRAGMENT).
GN PGI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=536;
RX MEDLINE=20351039; PubMed=10894541;
RA Reid S.D., Herbelin C.J., Bumbaugh A.C., Selander R.K., Whittam T.S.;
RT "Parallel evolution of virulence in pathogenic Escherichia coli.";
RL Nature 406:64-67(2000).
DR EMBL; AF267598; AAF97138.1; -.
DR InterPro; IPR001672; G6P_Isomerase.
DR Pfam; PF00342; PGI; 1.
DR PRINTS; PR00662; G6PISOMERASE.
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DR PROSITE; PS00765; P_GLUCOSE_ISOMERASE_1; 1.
KW Isomerase.
FT NON_TER 1
FT NON_TER 326
SQ SEQUENCE 326 AA; 36340 MW; C76930B3FOCF625A CRC64;

Query Match 69.1%; Score 38; DB 2; Length 326;
Best Local Similarity 66.7%; Pred. No. 13;
Matches 8; Conservative 2; Mismatches 0; Indels 0; Gaps 1;

QY 1 HHQKLV--FFAE 10
    |||||: |||
Db 289 HHQKLLSKFFAQ 300

RESULT 38
Q9SN52 PRELIMINARY; PRT; 191 AA.
AC Q9SN52;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DT 01-OCT-2000 (TREMBlrel. 15, Last annotation update)
DE HYPOTHETICAL 21.7 KDA PROTEIN.
GN F28A21.20 OR AT4G18610.
OS Arabidopsis thaliana (Mouse-ear cress).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
OC eurosids II; Brassicales; Brassicaceae; Arabidopsis.
OX NCBI_TaxID=3702;
RN [1]
RP SEQUENCE FROM N.A.
RA Bevan M., Mueller M.W., Muendlein A., Felber R., Bancroft I.,
RA Mewes H.W., Mayer K.F.X., Lemcke K., Schueller C.;
RL Submitted (FEB-1999) to the EMBL/GenBank/DBJ databases.
RN [2]
RP SEQUENCE FROM N.A.
RA EU Arabidopsis sequencing project;
RL Submitted (SEP-1999) to the EMBL/GenBank/DBJ databases.
RN [3]
RP SEQUENCE FROM N.A.
RA Mueller M.W., Muendlein A., Felber R., Mewes H.W., Lemcke K.,
RA Mayer K.F.X.;
RL Submitted (MAR-2000) to the EMBL/GenBank/DBJ databases.
RN [4]
RP SEQUENCE FROM N.A.
RA EU Arabidopsis sequencing project;
RL Submitted (MAR-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL; AL035526; CAB37446.1; -
DR EMBL; AL161549; CAB78863.1; -
KW Hypothetical protein.
SQ SEQUENCE 191 AA; 21744 MW; DFB6D3495AEB132F CRC64;

Query Match 67.3%; Score 37; DB 10; Length 191;
Best Local Similarity 60.0%; Pred. No. 12;
Matches 6; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
    ||| |||
Db 86 HHQACVFFGQ 95

RESULT 39
Q9U0M8 PRELIMINARY; PRT; 584 AA.
AC Q9U0M8;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE HYPOTHETICAL 71.0 KDA PROTEIN.
GN MALIP3.09.
OS Plasmodium falciparum (isolate 3D7).
OC Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.
OX NCBI_TaxID=36329;

[1]
RN RP SEQUENCE FROM N.A.
RC STRAIN-3D7; Churcher C., Harris B., Harris D., Lawson D., Quail M.,
RA Bowman S., Churcher C., Harris B., Harris D., Lawson D., Quail M.,
RA Barrell B.;
RL Submitted (SEP-1998) to the EMBL/GenBank/DBJ databases.
DR EMBL; AL031746; CAB63564.1; -
KW Hypothetical protein.
SQ SEQUENCE 584 AA; 70984 MW; 6E06F4C58A08F838 CRC64;

Query Match 67.3%; Score 37; DB 5; Length 584;
Best Local Similarity 50.0%; Pred. No. 38;
Matches 5; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
    |||||: |||
Db 528 HHQKTMFTQ 537

RESULT 40
Q93E19 PRELIMINARY; PRT; 1035 AA.
ID Q93E19
AC Q93E19;
DT 01-DEC-2001 (TREMBlrel. 19, Created)
DT 01-DEC-2001 (TREMBlrel. 19, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE ADEB RND PROTEIN.
GN ADEB.
OS Acinetobacter baumannii.
OC Bacteria; Proteobacteria; gamma subdivision; Moraxellaceae;
OC Acinetobacter.
OX NCBI_TaxID=470;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN-BM4454;
RA Magnet S., Courvalin P., Lambert T.;
RT "Characterization of a RND type efflux pump involved in aminoglycoside
RT resistance in Acinetobacter baumannii clinical isolate.";
RL Submitted (APR-2001) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF370885; AAL1440.1; -
SQ SEQUENCE 1035 AA; 112614 MW; 928E7935D84BFCF3 CRC64;

Query Match 67.3%; Score 37; DB 2; Length 1035;
Best Local Similarity 77.8%; Pred. No. 68;
Matches 7; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKLVFFFA 9
    |||| |||
Db 503 HHQKKGFFA 511

RESULT 41
Q9XST6 PRELIMINARY; PRT; 103 AA.
ID Q9XST6
AC Q9XST6;
DT 01-NOV-1999 (TREMBlrel. 12, Created)
DT 01-NOV-1999 (TREMBlrel. 12, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE TRANSMEMBRANE PROTEIN (FRAGMENT).
GN SAS.
OS Canis familiaris (Dog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis.
OX NCBI_TaxID=9615;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE-THYROID;
RX MEDLINE=20422104; PubMed=10964405;
RA Pichon B., Mercan D., Pouillon V., Christophe-Hobertus C.,
RA Christophe D.;
RT "A method for the large-scale cloning of nuclear proteins and nuclear
RT targeting sequences on a functional basis.";
RL Anal. Biochem. 284:231-239(2000).
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DR EMBL; AJ388529; CAB46831.1; -.
DR InterPro; IPR000301; Transmem_4.
DR Pfam; PF00335; transmembrane4; 1.
DR PRINTS; PR00259; TMFOUR.
FT NON_TER 103 103
SQ SEQUENCE 103 AA; 10723 MW; 5528A76F35FAC581 CRC64;

  Query Match      65.5%; Score 36; DB 6; Length 103;
  Best Local Similarity 75.0%; Pred. No. 11;
  Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 42
Q9CUY7 PRELIMINARY; PRT; 152 AA.
ID Q9CUY7
AC Q9CUY7
DT 01-JUN-2001 (TREMBlrel. 17, Created)
DT 01-JUN-2001 (TREMBlrel. 17, Last sequence update)
DT 01-JUN-2001 (TREMBlrel. 17, Last annotation update)
DE 2700085A14RIK PROTEIN (FRAGMENT).
GN 2700085A14RIK.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Euthera; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BL/6J; TISSUE=EMBRYO;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA Saito T., Okazaki Y., Gojohori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai H., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyooka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK012853; BAB28514.1; -.
DR EMBL; AK012567; BAB28322.1; -.
DR MGD; MGI:1914375; 2700085A14RIK.
DR InterPro; IPR000301; Transmem_4.
DR PRINTS; PR00259; TMFOUR.
FT NON_TER 152 152
SQ SEQUENCE 152 AA; 16162 MW; 5815EAA2F83F1B6D CRC64;

  Query Match      65.5%; Score 36; DB 11; Length 152;
  Best Local Similarity 75.0%; Pred. No. 16;
  Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 43
Q9CPW6 PRELIMINARY; PRT; 190 AA.
ID Q9CPW6
AC Q9CPW6

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DT 01-JUN-2001 (TREMBlrel. 17, Created)
DT 01-JUN-2001 (TREMBlrel. 17, Last sequence update)
DT 01-JUN-2001 (TREMBlrel. 17, Last annotation update)
DE 2700085A14RIK PROTEIN.
GN 2700085A14RIK.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Euthera; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BL/6J; TISSUE=EMBRYO;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA Saito T., Okazaki Y., Gojohori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai H., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyooka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK012853; BAB28514.1; -.
DR EMBL; AK012567; BAB28322.1; -.
DR MGD; MGI:1914375; 2700085A14RIK.
DR InterPro; IPR000301; Transmem_4.
DR PRINTS; PR00259; TMFOUR.
SQ SEQUENCE 190 AA; 20620 MW; EFB9D78DACD6927 CRC64;

  Query Match      65.5%; Score 36; DB 11; Length 190;
  Best Local Similarity 75.0%; Pred. No. 19;
  Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHQKLVFF 8
Db 50 HHQVLLFF 57

RESULT 44
Q9D8C2 PRELIMINARY; PRT; 204 AA.
ID Q9D8C2
AC Q9D8C2
DT 01-JUN-2001 (TREMBlrel. 17, Created)
DT 01-JUN-2001 (TREMBlrel. 17, Last sequence update)
DT 01-JUN-2001 (TREMBlrel. 17, Last annotation update)
DE 1100001123RIK PROTEIN.
GN 1100001123RIK.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Euthera; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BL/6J; TISSUE=SMALL INTESTINE;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA Saito T., Okazaki Y., Gojohori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai H., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyooka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK012853; BAB28514.1; -.
DR EMBL; AK012567; BAB28322.1; -.
DR MGD; MGI:1914375; 2700085A14RIK.
DR InterPro; IPR000301; Transmem_4.
DR PRINTS; PR00259; TMFOUR.
SQ SEQUENCE 190 AA; 20620 MW; EFB9D78DACD6927 CRC64;

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RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyo-oka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK0081175; BAB25510.1; -.
DR MGD; MGI:1913359; 1100001123Rik.
DR InterPro; IPR000301; Transmem_4.
DR Pfam; PF00335; transmembrane4; 1.
DR PRINTS; PR00259; TMFOUR.
SQ SEQUENCE 204 AA; 22219 MW; 76B95421EBCAE5F0 CRC64;

Query Match 65.5%; Score 36; DB 11; Length 204;
Best Local Similarity 75.0%; Pred. No. 21;
Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 45
Q9CQ88 PRELIMINARY; PRT; 210 AA.
AC Q9CQ88;
DT 01-JUN-2001 (TrEMBLrel. 17, Created)
DT 01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
DE 2700085A14RIK PROTEIN.
GN 2700085A14RIK.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BL/6J; TISSUE=EMBRYO;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA Saito T., Okazaki Y., Gojobori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Kuehl P., Lewis S., Matsuo Y., Nikaide I., Pesole G., Quackenbush J.,
RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyo-oka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK012571; BAB28326.1; -.
DR MGD; MGI:1913359; 1100001123Rik.
DR InterPro; IPR000301; Transmem_4.
DR Pfam; PF00335; transmembrane4; 1.
DR PRINTS; PR00259; TMFOUR.
SQ SEQUENCE 226 AA; 24566 MW; 684BAC91D7C42DEE CRC64;

Query Match 65.5%; Score 36; DB 11; Length 226;
Best Local Similarity 75.0%; Pred. No. 23;
Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 47
Q9UL10 PRELIMINARY; PRT; 396 AA.
AC Q9UL10;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DE 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE HYPOTHETICAL 42.4 KDA PROTEIN.
GN SARH.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

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Query Match 65.5%; Score 36; DB 11; Length 210;
Best Local Similarity 75.0%; Pred. No. 22;
Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 46
Q9CZ16 PRELIMINARY; PRT; 226 AA.
AC Q9CZ16;
DT 01-JUN-2001 (TrEMBLrel. 17, Created)
DT 01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
DT 01-JUN-2001 (TrEMBLrel. 17, Last annotation update)
DE 1100001123RIK PROTEIN.
GN 1100001123RIK.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BL/6J; TISSUE=EMBRYO;
RX MEDLINE=21085660; PubMed=11217851;
RA Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA Saito T., Okazaki Y., Gojobori T., Bono H., Kasukawa T., Saito R.,
RA Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA Kuehl P., Lewis S., Matsuo Y., Nikaide I., Pesole G., Quackenbush J.,
RA Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
RA Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA Suzuki H., Toyo-oka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.;
RT "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK012571; BAB28326.1; -.
DR MGD; MGI:1913359; 1100001123Rik.
DR InterPro; IPR000301; Transmem_4.
DR Pfam; PF00335; transmembrane4; 1.
DR PRINTS; PR00259; TMFOUR.
SQ SEQUENCE 226 AA; 24566 MW; 684BAC91D7C42DEE CRC64;

Query Match 65.5%; Score 36; DB 11; Length 226;
Best Local Similarity 75.0%; Pred. No. 23;
Matches 6; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQKLVFF 8
Db 70 HHQVLLFF 77

RESULT 47
Q9UL10 PRELIMINARY; PRT; 396 AA.
AC Q9UL10;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DE 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE HYPOTHETICAL 42.4 KDA PROTEIN.
GN SARH.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

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OC Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=LIVER;
RX MEDLINE=99375323; PubMed=10444331;
RA Eschenbrenner M., Schuman Joris M.;
RT "Cloning and mapping of the cDNA for human sarcosine dehydrogenase, a
RL flavoenzyme defective in patients with sarcosinemia.";
RL Genomics 59:300-308(1999).
DR EMBL: AF095737; AAD53400.2; -.
DR InterPro: IPR000527; DAO.
DR InterPro: IPR000205; NAD_binding.
DR Pfam: PF01266; DAO; 1.
KW Hypothetical protein.
SQ SEQUENCE 396 AA; 42362 MW; 150CA3706476BB69 CRC64;

Query Match 65.5%; Score 36; DB 4; Length 396;
Best Local Similarity 62.5%; Pred. NO. 41;
Matches 5; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQKLVFF 8
Db 348 HHTRLIFF 355

RESULT 48
Q01165 PRELIMINARY; PRT; 535 AA.
AC Q01165;
DT 01-NOV-1996 (TRENBLrel. 01, Created)
DT 01-NOV-1996 (TRENBLrel. 01, Last sequence update)
DT 01-MAR-2001 (TRENBLrel. 16, Last annotation update)
DE TRANSPOSASE.
OS Magnaporthe grisea (Rice blast fungus) (Pyricularia grisea).
OC Eukaryota; Fungi; Ascomycota; Pezizomycotina; Sordariomycetes;
OC Sordariomycetes incertae sedis; Magnaporthaceae; Magnaporthe.
OX NCBI_TaxID=148305;
RN [1]
RP SEQUENCE FROM N.A.
RC TRANSPOSON-POT2;
RX MEDLINE=95115685; PubMed=7816044;
RA Kachroo P., Leong S.A., Chaitoo B.;
RT "Pot2, an inverted repeat transposon from the rice blast fungus
RL Magnaporthe grisea.";
RL Mol. Gen. Genet. 245:339-348(1994).
DR EMBL: Z33638; CAA83918.1; -.
SQ SEQUENCE 535 AA; 61079 MW; A755F73FE6878F47 CRC64;

Query Match 65.5%; Score 36; DB 3; Length 535;
Best Local Similarity 77.8%; Pred. NO. 55;
Matches 7; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HOKLVFFAE 10
Db 80 HOELRFFAE 88

RESULT 49
Q26556 PRELIMINARY; PRT; 859 AA.
ID Q26556;
AC Q26556;
DT 01-JAN-1998 (TRENBLrel. 05, Created)
DT 01-JAN-1998 (TRENBLrel. 05, Last sequence update)
DT 01-OCT-2001 (TRENBLrel. 18, Last annotation update)
DE MAGNESIUM CHELATASE SUBUNIT.
GN MTH456.
OS Methanothermobacter thermoautotrophicus.
OC Archaea; Euryarchaeota; Methanobacteriales; Methanobacteriaceae;
OC Methanothermobacter.
OX NCBI_TaxID=145262;
RN [1]
RP SEQUENCE FROM N.A.
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RC STRAIN-DELTA H;
RX MEDLINE=98037514; PubMed=9371463;
RA Smith D.R., Doucette-Stamm L.A., DeLoughery C., Lee H.-M., DuBois J.,
RA Aldredge T., Bashirzadeh R., Blakely D., Cook R., Gilbert K.,
RA Harrison D., Hoang L., Keagle P., Lum W., Pothier B., Qiu D.,
RA Spadafora R., Vicare R., Wang Y., Wierzbowski J., Gibson R.,
RA Jiwani N., Caruso A., Bush D., Safer H., Patwell D., Prabhakar S.,
RA McDougall S., Shimer G., Goyal A., Pietrovski S., Church G.M.,
RA Daniels C.J., Mao J.-I., Rice P., Nolling J., Reeve J.N.;
RT "Complete genome sequence of Methanobacterium thermoautotrophicum
RT deltaH: functional analysis and comparative genomics.";
RL J. Bacteriol. 179:7135-7155(1997).
RL EMBL: AE000868; AAB85426.1; -.
DR InterPro: IPR003672; COBN/Mg_chelataase.
DR Pfam: PF02514; COBN-Mg_chel. 1.
KW Complete proteome.
SQ SEQUENCE 859 AA; 97572 MW; 0C7946B7839EF5ED CRC64;

Query Match 65.5%; Score 36; DB 17; Length 859;
Best Local Similarity 66.7%; Pred. NO. 89;
Matches 6; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

Qy 1 HHQKLVFFA 9
Db 210 HHQYLAFYA 218

RESULT 50
Q27011 PRELIMINARY; PRT; 1668 AA.
ID Q27011;
AC Q27011;
DT 01-JAN-1998 (TRENBLrel. 05, Created)
DT 01-JAN-1998 (TRENBLrel. 05, Last sequence update)
DT 01-OCT-2001 (TRENBLrel. 18, Last annotation update)
DE COBALAMIN BIOSYNTHESIS PROTEIN N.
GN MTH928.
OS Methanothermobacter thermoautotrophicus.
OC Archaea; Euryarchaeota; Methanobacteriales; Methanobacteriaceae;
OC Methanothermobacter.
OX NCBI_TaxID=145262;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN-DELTA H;
RX MEDLINE=98037514; PubMed=9371463;
RA Smith D.R., Doucette-Stamm L.A., DeLoughery C., Lee H.-M., DuBois J.,
RA Aldredge T., Bashirzadeh R., Blakely D., Cook R., Gilbert K.,
RA Harrison D., Hoang L., Keagle P., Lum W., Pothier B., Qiu D.,
RA Spadafora R., Vicare R., Wang Y., Wierzbowski J., Gibson R.,
RA Jiwani N., Caruso A., Bush D., Safer H., Patwell D., Prabhakar S.,
RA McDougall S., Shimer G., Goyal A., Pietrovski S., Church G.M.,
RA Daniels C.J., Mao J.-I., Rice P., Nolling J., Reeve J.N.;
RT "Complete genome sequence of Methanobacterium thermoautotrophicum
RT deltaH: functional analysis and comparative genomics.";
RL J. Bacteriol. 179:7135-7155(1997).
RL EMBL: AE000868; AAB85426.1; -.
DR InterPro: IPR003672; COBN/Mg_chelataase.
DR InterPro: IPR001993; Mitoch_carrier.
DR Pfam: PF02514; COBN-Mg_chel. 1.
DR PROSITE: PS00215; MITOCH_CARRIER; UNKNOWN_1.
KW Complete proteome.
SQ SEQUENCE 1668 AA; 184731 MW; 73D53E89519EAC00 CRC64;

Query Match 65.5%; Score 36; DB 17; Length 1668;
Best Local Similarity 66.7%; Pred. NO. 1.7e+02;
Matches 6; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

Qy 1 HHQKLVFFA 9
Db 792 HHQYLAFYA 800

Search completed: October 29, 2002, 09:25:20
Job time : 28 secs
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GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:37:12 : Search time 13 Seconds
(without alignments)
18.789 Million cell updates/sec

Title: US-09-724-842A-27

Perfect score: 55

Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 231628 seqs, 24425594 residues

Total number of hits satisfying chosen parameters: 70601

Minimum DB seq length: 0

Maximum DB seq length: 10

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 50 summaries

Database : Issued_Patents_AA:*

1: /cgn2_6/ptodata/1/iaa/5A_COMB.pep.*

2: /cgn2_6/ptodata/1/iaa/5B_COMB.pep.*

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6: /cgn2_6/ptodata/1/iaa/backfiles1.pep.*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	46	83.6	9	4	US-09-264-709A-4
2	42	76.4	8	2	US-08-612-785B-5
3	42	76.4	8	4	US-08-703-675C-28
4	42	76.4	8	4	US-08-617-267C-5
5	38	69.1	7	2	US-08-612-785B-6
6	38	69.1	7	4	US-08-703-675C-29
7	38	69.1	7	4	US-08-617-267C-6
8	34	61.8	7	1	US-08-127-904-14
9	34	61.8	7	1	US-08-397-633A-105
10	34	61.8	7	2	US-08-612-785B-7
11	34	61.8	7	4	US-08-703-675C-30
12	34	61.8	7	4	US-08-617-267C-7
13	34	61.8	7	4	US-09-264-709A-13
14	34	61.8	7	5	PCT-US94-10475-14
15	34	61.8	8	2	US-08-630-645-1
16	34	61.8	8	5	PCT-US96-10220-1
17	34	61.8	10	3	US-08-970-833-3
18	30	54.5	6	2	US-08-612-785B-8
19	30	54.5	6	2	US-08-461-216-3
20	30	54.5	6	4	US-08-703-675C-31
21	30	54.5	6	4	US-09-242-724-24
22	30	54.5	6	4	US-08-617-267C-8
23	30	54.5	6	4	US-08-723-661B-3
24	29	52.7	6	2	US-08-612-785B-9
25	29	52.7	6	2	US-08-612-785B-27
26	29	52.7	6	4	US-08-703-675C-32
27	29	52.7	6	4	US-08-703-675C-40

ALIGNMENTS

RESULT 1

US-09-264-709A-4

; Sequence 4, Application US/09264709A

; Patent No. 6320024

; GENERAL INFORMATION:

; APPLICANT: Roberts, Eugene

; TITLE OF INVENTION: Method for Design of Substances that Enhance Memory and

; TITLE OF INVENTION: Improve the Quality of Life

; FILE REFERENCE: 2124-310

; CURRENT APPLICATION NUMBER: US/09/264,709A

; CURRENT FILING DATE: 1999-03-09

; PRIOR APPLICATION NUMBER: 08/797,782

; PRIOR FILING DATE: 1997-02-07

; NUMBER OF SEQ ID NOS: 39

; SOFTWARE: PatentIn Ver. 2.1

; SEQ ID NO 4

; LENGTH: 9

; TYPE: PRT

; ORGANISM: Homo sapiens

US-09-264-709A-4

Query Match 83.6%; Score 46; DB 4; Length 9;

Best Local Similarity 100.0%; Pred. No. 1.7e+05;

Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 HHQKLVFF 8

Db 2 HHQKLVFF 9

|||||

RESULT 2

US-08-612-785B-5

; Sequence 5, Application US/08612785B

; Patent No. 5854204

; GENERAL INFORMATION:

; APPLICANT: Findeis, Mark A. et al.

; TITLE OF INVENTION: AB Peptides that Modulate b-Amyloid

; TITLE OF INVENTION: Aggregation

; NUMBER OF SEQUENCES: 40

; CORRESPONDENCE ADDRESS:

; ADDRESSEE: LAHIVE & COCKFIELD

; STREET: 28 State Street, Suite 510

; CITY: Boston

; STATE: Massachusetts

; COUNTRY: USA

Sequence 9, Appli
Sequence 27, Appli
Sequence 27, Appli
Sequence 30, Appli
Sequence 1, Appli
Sequence 33, Appli
Sequence 15, Appli
Sequence 10, Appli
Sequence 2, Appli
Sequence 46, Appli
Sequence 25, Appli
Sequence 26, Appli
Sequence 10, Appli
Sequence 28, Appli
Sequence 15, Appli
Sequence 31, Appli
Sequence 44, Appli
Sequence 31, Appli
Sequence 43, Appli
Sequence 4, Appli
Sequence 11, Appli

ZIP: 02109-1875
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/612,785B
FILING DATE: Herewith
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 5:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-612-785B-5

Query Match 76.4%; Score 42; DB 2; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2 HQKLVFFA 9
Db 1 HQKLVFFA 8
|||||||

RESULT 3
US-08-703-675C-28
Sequence 28, Application US/08703675C
Patent No. 6303567
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Modulators of -Amyloid Peptide
NUMBER OF SEQUENCES: 46
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/703,675C
FILING DATE: 27-AUG-1996
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995

Aggregation Comprising D-

PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/616,081
FILING DATE: 14-MAR-1996
ATTORNEY/AGENT INFORMATION:
NAME: Kara, Catherine J.
REGISTRATION NUMBER: 41,106
REFERENCE/DOCKET NUMBER: PPI-016CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 28:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-703-675C-28

Query Match 76.4%; Score 42; DB 4; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2 HQKLVFFA 9
Db 1 HQKLVFFA 8
|||||||

RESULT 4
US-08-617-267C-5
Sequence 5, Application US/08617267C
Patent No. 6319498
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Modulators of Amyloid Aggregation
NUMBER OF SEQUENCES: 45
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/617,267C
FILING DATE: 14-MAR-1996
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)227-5941
INFORMATION FOR SEQ ID NO: 5:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
TYPE: amino acid

; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-617-267C-5

Query Match 76.4%; Score 42; DB 4; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HOKLVFFA 9
| | | | |
Db 1 HOKLVFFA 8

RESULT 5

US-08-612-785B-6
; Sequence 6, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: AB Peptides that Modulate b-Amyloid
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/612.785B
FILING DATE: Herewith
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
LENGTH: 7 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide

US-08-612-785B-6

Query Match 69.1%; Score 38; DB 2; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HOKLVFF 8
| | | | |
Db 1 HOKLVFF 7

RESULT 6

US-08-703-675C-29

; Sequence 29, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/703.675C
FILING DATE: 27-AUG-1996
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/616,081
FILING DATE: 14-MAR-1996
ATTORNEY/AGENT INFORMATION:
NAME: Kara, Catherine J.
REGISTRATION NUMBER: 41,106
REFERENCE/DOCKET NUMBER: PPI-016CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 29:
SEQUENCE CHARACTERISTICS:
LENGTH: 7 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-703-675C-29

Query Match 69.1%; Score 38; DB 4; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HOKLVFF 8
| | | | |
Db 1 HOKLVFF 7

RESULT 7

US-08-617-267C-6
; Sequence 6, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/617,267C
FILING DATE: 14-MAR-1996
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP2
TELEPHONE: (617)227-7400
TELEFAX: (617)227-5941
INFORMATION FOR SEQ ID NO: 6:
SEQUENCE CHARACTERISTICS:
LENGTH: 7 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-617-267C-6

Query Match 69.1%; Score 38; DB 4; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0;

QY 2 HQKLVFF 8
DB 1 HQKLVFF 7

RESULT 8
US-08-127-904-14
Sequence 14, Application US/08127904
Patent No. 5470951
GENERAL INFORMATION:
APPLICANT: Eugene Roberts
TITLE OF INVENTION: Method For Antagonizing
TITLE OF INVENTION: Amnestic Effects of Amyloid n
TITLE OF INVENTION: Protein and Improving the
TITLE OF INVENTION: Quality of Life in Individuals
TITLE OF INVENTION: With Alzheimer Disease
NUMBER OF SEQUENCES: 15
CORRESPONDENCE ADDRESS:
ADDRESSEE: City of Hope
STREET: 1500 East Duarte Road
CITY: Duarte
STATE: California
COUNTRY: United States of America
ZIP: 91010-0269
COMPUTER READABLE FORM:
MEDIUM TYPE: 3M Double Density 5 1/4" diskette
COMPUTER: Wang PC
OPERATING SYSTEM: MS DOS Version 3.20
SOFTWARE: Microsoft
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/127,904
FILING DATE: 29 September 1993
CLASSIFICATION: 424
PRIOR APPLICATION DATA: NO. 5470951e
ATTORNEY/AGENT INFORMATION:
NAME: Irons, Edward S.
REGISTRATION NUMBER: 16,541

REFERENCE/DOCKET NUMBER: No. 5470951e
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 783-6040
TELEFAX: (202) 783-6031
TELEX: No. 5470951e
INFORMATION FOR SEQ ID NO: 14:
SEQUENCE CHARACTERISTICS:
LENGTH: 7
TYPE: Amino Acid
STRANDEDNESS:
TOPOLOGY: Unknown
US-08-127-904-14

Query Match 61.8%; Score 34; DB 1; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0;

QY 4 KLVFFAE 10
DB 1 KLVFFAE 7

RESULT 9
US-08-397-633A-105
Sequence 105, Application US/08397633A
Patent No. 5773577
GENERAL INFORMATION:
APPLICANT: Cappello, Joseph
TITLE OF INVENTION: PRODUCTS COMPRISING SUBSTRATESCAPABLE
TITLE OF INVENTION: OF ENZYMAIC CROSS-LINKING
NUMBER OF SEQUENCES: 105
CORRESPONDENCE ADDRESS:
ADDRESSEE: FLEHR, HOHBACH, TEST, ALBRITTON & HERBERT
STREET: 4 Embarcadero Center, Suite 3400
CITY: San Francisco
STATE: California
COUNTRY: USA
ZIP: 94111-4187
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/397,633A
FILING DATE:
CLASSIFICATION: 530
ATTORNEY/AGENT INFORMATION:
NAME: Rowland, Bertram I.
REGISTRATION NUMBER: 20,015
REFERENCE/DOCKET NUMBER: A-58848-1/BIR PROP-011-1
TELECOMMUNICATION INFORMATION:
TELEPHONE: (415) 781-1989
TELEFAX: (415) 398-3249
TELEX: 910 277299
INFORMATION FOR SEQ ID NO: 105:
SEQUENCE CHARACTERISTICS:
LENGTH: 7 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-397-633A-105

Query Match 61.8%; Score 34; DB 1; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0;

QY 1 HQKLV 6
DB 2 HQKLV 7

```
RESULT 10
US-08-612-785B-7
; Sequence 7, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Fideis, Mark A. et al.
; TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USN 08/404,831
; FILING DATE: 14-MAR-1995
; APPLICATION TYPE: peptide
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 7:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 7 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-612-785B-7

Query Match 61.8%; Score 34; DB 2; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFFA 9
Db 1 QKLVFFA 7

RESULT 11
US-08-703-675C-30
; Sequence 30, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Fideis, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USN 08/404,831
; FILING DATE: 14-MAR-1995
; APPLICATION TYPE: peptide
; ATTORNEY/AGENT INFORMATION:
; NAME: Kara, Catherine J.
; REGISTRATION NUMBER: 41,106
; REFERENCE/DOCKET NUMBER: PPI-016CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 30:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 7 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-703-675C-30

Query Match 61.8%; Score 34; DB 4; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFFA 9
Db 1 QKLVFFA 7

RESULT 12
US-08-617-267C-7
; Sequence 7, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:
; APPLICANT: Fideis, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USN 08/404,831
; FILING DATE: 14-MAR-1995
; APPLICATION TYPE: peptide
; ATTORNEY/AGENT INFORMATION:
; NAME: Kara, Catherine J.
; REGISTRATION NUMBER: 41,106
; REFERENCE/DOCKET NUMBER: PPI-016CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 30:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 7 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-703-675C-30
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; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-0020CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 7:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 7 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-617-267C-7

Query Match 61.8%; Score 34; DB 4; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFFA 9
Db 1 QKLVFFA 7

RESULT 13
US-09-264-709A-13
; Sequence 13, Application US/09264709A
; Patent No. 6320024
; GENERAL INFORMATION:
; APPLICANT: Roberts, Eugene
; TITLE OF INVENTION: Method for Design of Substances that Enhance Memory and
; TITLE OF INVENTION: Improve the Quality of Life
; FILE REFERENCE: 2124-310
; CURRENT APPLICATION NUMBER: US/09/264,709A
; CURRENT FILING DATE: 1999-03-09
; PRIOR APPLICATION NUMBER: 08/797,782
; PRIOR FILING DATE: 1997-02-07
; NUMBER OF SEQ ID NOS: 39
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 13
; LENGTH: 7
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-264-709A-13

Query Match 61.8%; Score 34; DB 4; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
Db 1 KLVFFAE 7

RESULT 14
PCT-US94-10475-14
; Sequence 14, Application PC/TUS9410475
; GENERAL INFORMATION:
; APPLICANT: Eugene Roberts
; TITLE OF INVENTION: Method For
; TITLE OF INVENTION: Antagonizing Amnestic
; TITLE OF INVENTION: Effects of Amyloid n
; TITLE OF INVENTION: Protein and Improving
; TITLE OF INVENTION: the Quality of Life
; TITLE OF INVENTION: in Individuals
; TITLE OF INVENTION: With Alzheimer Disease
; NUMBER OF SEQUENCES: 15
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: City of Hope
;
```

```
; STREET: 1500 East Duarte Road
; CITY: Duarte
; STATE: California
; COUNTRY: United States of America
; ZIP: 91010-0269
; COMPUTER READABLE FORM:
; MEDIUM TYPE: 3M Double Density 5 1/4"
; MEDIUM TYPE: diskette
; COMPUTER: Wang PC
; OPERATING SYSTEM: MS DOS Version 3.20
; SOFTWARE: Microsoft
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: PCT/US94/10475
; FILING DATE: 16 September 1994
; CLASSIFICATION:
; PRIOR APPLICATION DATA: U. S. Application
; PRIOR APPLICATION DATA: Serial No.
; PRIOR APPLICATION DATA: 08/127,904; filed
; PRIOR APPLICATION DATA: 29 September 1993
; ATTORNEY/AGENT INFORMATION:
; NAME: Irons, Edward S.
; REGISTRATION NUMBER: 16,541
; REFERENCE/DOCKET NUMBER: None
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 626-3564 or 783-6030
; TELEFAX: (202) 783-6031
; TELEX: None
; INFORMATION FOR SEQ ID NO: 14:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 7
; TYPE: Amino Acid
; STRANDEDNESS:
; TOPOLOGY: Unknown
PCT-US94-10475-14

Query Match 61.8%; Score 34; DB 5; Length 7;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFAE 10
Db 1 KLVFFAE 7

RESULT 15
US-08-630-645-1
; Sequence 1, Application US/08630645
; Patent No. 5948763
; GENERAL INFORMATION:
; APPLICANT: SOTO-JARA, Claudio
; APPLICANT: BAUMANN, Marc
; APPLICANT: FRANGIONE, Blas
; TITLE OF INVENTION: PEPTIDES AND PHARMACEUTICAL COMPOSITIONS
; TITLE OF INVENTION: THEREOF FOR TREATMENT OF DISORDERS OR DISEASES ASSOCIATED
; TITLE OF INVENTION: WITH PROTEIN FOLDING INTO AMYLOID OR AMYLOID-LIKE DEPOSITS
; NUMBER OF SEQUENCES: 26
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: BROWDY AND NEIMARK
; STREET: 419 Seventh Street, N.W., Suite 400
; CITY: Washington
; STATE: D.C.
; COUNTRY: USA
; ZIP: 20004
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/630,645
; FILING DATE:
; CLASSIFICATION: 530
; PRIOR APPLICATION DATA:
;
```


APPLICATION NUMBER: US 08/478,326
FILING DATE: 06-JUN-1995
ATTORNEY/AGENT INFORMATION:
NAME: YUN, Allen C.
REGISTRATION NUMBER: 37,971
REFERENCE/DOCKET NUMBER: SOTO-JARA-1
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-628-5197
TELEFAX: 202-737-3528
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-630-645-1

Query Match 61.8%; Score 34; DB 2; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFFAE 10
Db 1 KLVFFAE 7

RESULT 16

PCT-US96-10220-1
Sequence 1, Application PC/TUS9610220
GENERAL INFORMATION:
APPLICANT:
TITLE OF INVENTION: PEPTIDES AND PHARMACEUTICAL COMPOSITIONS
TITLE OF INVENTION: THEROF FOR TREATMENT OF DISORDERS OR DISEASES ASSOCIATED
TITLE OF INVENTION: WITH PROTEIN FOLDING INTO AMYLOID OR AMYLOID-LIKE DEPOSITS
NUMBER OF SEQUENCES: 26
CORRESPONDENCE ADDRESS:
ADDRESS: BROWDY AND NEIMARK
STREET: 419 Seventh Street, N.W., Suite 400
CITY: Washington
STATE: D.C.
COUNTRY: USA
ZIP: 20004
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US96/10220
FILING DATE:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/478,326
FILING DATE: 06-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/630,645
FILING DATE: 10-APR-1996
ATTORNEY/AGENT INFORMATION:
NAME: BROWDY, Roger L.
REGISTRATION NUMBER: 25,618
REFERENCE/DOCKET NUMBER: SOTO-JARA-1 PCT
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-628-5197
TELEFAX: 202-737-3528
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 8 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
PCT-US96-10220-1

Query Match 61.8%; Score 34; DB 5; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFFAE 10
Db 1 KLVFFAE 7

RESULT 17

US-08-970-833-3
Sequence 3, Application US/08970833
Patent No. 602859
GENERAL INFORMATION:
APPLICANT: Kiessling, Laura L.
APPLICANT: Murphy, Regina M.
TITLE OF INVENTION: INHIBITORS OF BETA-AMYLOID TOXICITY
NUMBER OF SEQUENCES: 11
CORRESPONDENCE ADDRESS:
ADDRESS: Quarles & Brady
STREET: 411 East Wisconsin Avenue
CITY: Milwaukee
STATE: Wisconsin
COUNTRY: U.S.A.
ZIP: 53202-4497
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/970,833
FILING DATE:
CLASSIFICATION: 530
ATTORNEY/AGENT INFORMATION:
NAME: Baker, Jean C.
REGISTRATION NUMBER: 35,433
REFERENCE/DOCKET NUMBER: 960296.94291
TELECOMMUNICATION INFORMATION:
TELEPHONE: (414) 277-5709
TELEFAX: (414) 271-3552
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
LENGTH: 10 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-970-833-3

Query Match 61.8%; Score 34; DB 3; Length 10;
Best Local Similarity 100.0%; Pred. No. 0.89;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFFAE 10
Db 1 KLVFFAE 7

RESULT 18

US-08-612-785B-8
Sequence 8, Application US/08612785B
Patent No. 5854204
GENERAL INFORMATION:
APPLICANT: Findeis, Mark A. et al.
TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
TITLE OF INVENTION: Aggregation
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESS:
ADDRESS: LAHIVE & COCKFIELD
STREET: 28 State Street, Suite 510
CITY: Boston
STATE: Massachusetts

```

; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent in Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 8:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-612-785B-8

Query Match 54.5%; Score 30; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFF 8
Db 1 QKLVFF 6

RESULT 19
US-08-461-216-3
; Sequence 3, Application US/08461216
; Patent No. 5958883
; GENERAL INFORMATION:
; APPLICANT: Snow, A.D.
; TITLE OF INVENTION: ANIMAL MODELS OF HUMAN AMYLOIDOSES
; NUMBER OF SEQUENCES: 8
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Christensen, O'Connor, Johnson and Kindness
; STREET: 1420 Fifth Avenue, Suite 2800
; CITY: Seattle
; STATE: Washington
; COUNTRY: USA
; ZIP: 98101-2347
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette-5.25 inch, 1.2Mb storage
; COMPUTER: IBM PC/386 Compatible
; OPERATING SYSTEM: MS-DOS 4.01
; SOFTWARE: Word for Windows-t
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/461,216
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 07/969,734
; FILING DATE: October 23, 1992
; APPLICATION NUMBER: 07/950,417
; FILING DATE: September 23, 1992

```

```

; ATTORNEY/AGENT INFORMATION:
; NAME: Broderick, Thomas F.
; REGISTRATION NUMBER: 31,332
; REFERENCE/DOCKET NUMBER: UOFW-1-6707
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 1-206-682-8100; 1-206-224-0709 (direct)
; TELEFAX: 1-206-224-0779
; TELEX: 4938023
; INFORMATION FOR SEQ ID NO: 3:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; DESCRIPTION: [SYMBOL 98 \f "Symbol"]/M4(12-17);
; DESCRIPTION: page 60, line 4-5; page 83, line 33 and 27-28
US-08-461-216-3

Query Match 54.5%; Score 30; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HQOKL 5
Db 2 HQOKL 6

RESULT 20
US-08-703-675C-31
; Sequence 31, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent in Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/703,675C
; FILING DATE: 27-AUG-1996
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/616,081
; FILING DATE: 14-MAR-1996
; ATTORNEY/AGENT INFORMATION:
; NAME: Kara, Catherine J.
; REGISTRATION NUMBER: 41,106
; REFERENCE/DOCKET NUMBER: PPI-016CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 31:
; SEQUENCE CHARACTERISTICS:

```

Aggregation Comprising

; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-703-675C-31

Query Match 54.5%; Score 30; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QKLVEF 8
Db 1 QKLVEF 6

RESULT 21

US-09-242-724-24

; Sequence 24, Application US/09242724

; Patent No. 6316405

; GENERAL INFORMATION:

; APPLICANT: Solomon, Michael E.

; APPLICANT: Rich, Daniel H.

; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor

; FILE REFERENCE: Cyclosporin Analogs

; CURRENT APPLICATION NUMBER: US/09/242.724

; CURRENT FILING DATE: 1999-02-22

; NUMBER OF SEQ ID NOS: 33

; SOFTWARE: PatentIn Ver. 2.0

; SEQ ID NO 24

; LENGTH: 6

; TYPE: PRT

; ORGANISM: Artificial Sequence

; FEATURE:

; OTHER INFORMATION: ;

US-09-242-724-24

Query Match 54.5%; Score 30; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QKLVEF 8
Db 1 QKLVEF 6

RESULT 22

US-08-617-267C-8

; Sequence 8, Application US/08617267C

; Patent No. 6319498

; GENERAL INFORMATION:

; APPLICANT: Findels, Mark A. et al.

; TITLE OF INVENTION: Modulators of Amyloid Aggregation

; NUMBER OF SEQUENCES: 45

; CORRESPONDENCE ADDRESS:

; ADDRESSEE: LAHIVE & COCKFIELD, LLP

; STREET: 28 State Street

; CITY: Boston

; STATE: Massachusetts

; COUNTRY: USA

; ZIP: 02109-1875

; COMPUTER READABLE FORM:

; MEDIUM TYPE: Floppy disk

; COMPUTER: IBM PC compatible

; OPERATING SYSTEM: PC-DOS/MS-DOS

; SOFTWARE: PatentIn Release #1.0, Version #1.25

; CURRENT APPLICATION DATA:

; APPLICATION NUMBER: US/08/617.267C

; FILING DATE: 14-MAR-1996

; PRIOR APPLICATION DATA:

; APPLICATION NUMBER: USSN 08/404,831

; FILING DATE: 14-MAR-1995

; PRIOR APPLICATION DATA:

; APPLICATION NUMBER: USSN 08/475,579

; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 8:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-617-267C-8

Query Match 54.5%; Score 30; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QKLVEF 8
Db 1 QKLVEF 6

RESULT 23

US-08-723-661B-3

; Sequence 3, Application US/08723661B

; Patent No. 6340783

; GENERAL INFORMATION:

; APPLICANT: Alan D Snow

; TITLE OF INVENTION: Animal Models of Human Amyloidoses

; NUMBER OF SEQUENCES: 7

; CORRESPONDENCE ADDRESS:

; ADDRESSEE: Patrick M. Dwyer

; STREET: 1818 Westlake Avenue N, Suite 114

; CITY: Seattle

; STATE: WA (Washington)

; COUNTRY: United States of America

; ZIP: 98109

; COMPUTER READABLE FORM:

; MEDIUM TYPE: Diskette - 3.50 inch, 1.44 Mb storage

; COMPUTER: IBM PC

; OPERATING SYSTEM: PC-DOS (Windows 98)

; SOFTWARE: WordPerfect 5.2

; CURRENT APPLICATION DATA:

; APPLICATION NUMBER: US/08/723,661B

; FILING DATE: 31-Oct-1996

; PRIOR APPLICATION DATA:

; APPLICATION NUMBER: 08/461,216

; FILING DATE: 05-Jun-1995

; APPLICATION NUMBER: 07/969,734

; FILING DATE: 23-Oct-1992

; APPLICATION NUMBER: 07/950,417

; FILING DATE: 23-Sep-1992

; ATTORNEY/AGENT INFORMATION:

; NAME: Dwyer, Patrick M.

; REGISTRATION NUMBER: 32,411

; REFERENCE/DOCKET NUMBER: PROTEO.P00C1

; TELECOMMUNICATION INFORMATION:

; TELEPHONE: (206) 343-7074

; TELEFAX: (206) 343-7085

; INFORMATION FOR SEQ ID NO: 3:

; SEQUENCE CHARACTERISTICS:

; LENGTH: 6 AMINO ACIDS

; TYPE: AMINO ACID

; STRANDEDNESS: SINGLE

; TOPOLOGY: LINEAR

; MOLECULE TYPE: PEPTIDE

; DESCRIPTION: /A4 (12-17); page 60, lines 4-5; page 83,

; SEQUENCE DESCRIPTION: SEQ ID NO: 3;
US-08-723-661B-3

Query Match 54.5%; Score 30; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKL 5
Db 2 HHQKL 6

RESULT 24

US-08-612-785B-9
; Sequence 9, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 9:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-612-785B-9

Query Match 52.7%; Score 29; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFFA 9
Db 1 KLVFFA 6

RESULT 25

US-08-612-785B-27
; Sequence 27, Application US/08612785B

; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 27:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-612-785B-27

Query Match 52.7%; Score 29; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFFAE 10
Db 1 LVFFAE 6

RESULT 26
US-08-703-675C-32
; Sequence 32, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible

OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/703,675C
FILING DATE: 27-AUG-1996
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/616,081
FILING DATE: 14-MAR-1996
ATTORNEY/AGENT INFORMATION:
NAME: Kara, Catherine J.
REGISTRATION NUMBER: 41,106
REFERENCE/DOCKET NUMBER: PPI-016CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 32:
SEQUENCE CHARACTERISTICS:
LENGTH: 6 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-703-675C-32

Query Match 52.7%; Score 29; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0;

QY 4 KLVEFFA 9
| | | | |
DB 1 KLVEFFA 6

RESULT 27
US-08-703-675C-40
Sequence 40, Application US/08703675C
Patent No. 6303567
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Modulators of -Amyloid Peptide
NUMBER OF SEQUENCES: 46
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/703,675C
FILING DATE: 27-AUG-1996
CLASSIFICATION: 514
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:

APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/616,081
FILING DATE: 14-MAR-1996
ATTORNEY/AGENT INFORMATION:
NAME: Kara, Catherine J.
REGISTRATION NUMBER: 41,106
REFERENCE/DOCKET NUMBER: PPI-016CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)742-4214
INFORMATION FOR SEQ ID NO: 40:
SEQUENCE CHARACTERISTICS:
LENGTH: 6 amino acids
TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-08-703-675C-40

Query Match 52.7%; Score 29; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0;

QY 5 LVFFAE 10
| | | | |
DB 1 LVFFAE 6

RESULT 28
US-08-617-267C-9
Sequence 9, Application US/08617267C
Patent No. 6319498
GENERAL INFORMATION:
APPLICANT: Findels, Mark A. et al.
TITLE OF INVENTION: Modulators of Amyloid Aggregation
NUMBER OF SEQUENCES: 45
CORRESPONDENCE ADDRESS:
ADDRESSEE: LAHIVE & COCKFIELD, LLP
STREET: 28 State Street
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875
COMPUTER READABLE FORM:
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/617,267C
FILING DATE: 14-MAR-1996
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/404,831
FILING DATE: 14-MAR-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/475,579
FILING DATE: 07-JUN-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: USSN 08/548,998
FILING DATE: 27-OCT-1995
ATTORNEY/AGENT INFORMATION:
NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503
REFERENCE/DOCKET NUMBER: PPI-002CP2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)227-7400
TELEFAX: (617)227-5941
INFORMATION FOR SEQ ID NO: 9:
SEQUENCE CHARACTERISTICS:
LENGTH: 6 amino acids
TYPE: amino acid
TOPOLOGY: linear

; MOLECULE TYPE: peptide
US-08-617-267C-9

Query Match 52.7%; Score 29; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVEFA 9
| | | | |
DB 1 KLVEFA 6

RESULT 29

US-08-617-267C-27
; Sequence 27, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 27:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-617-267C-27

Query Match 52.7%; Score 29; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFFAE 10
| | | | |
DB 1 LVFFAE 6

RESULT 30

US-09-242-724-27
; Sequence 27, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:

; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 27
; LENGTH: 6
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: synthetic
; OTHER INFORMATION: polypeptide
; NAME/KEY: MOD_RES
; LOCATION: (1)
; OTHER INFORMATION: ACETYLATION
; NAME/KEY: MOD_RES
; LOCATION: (2)
; OTHER INFORMATION: K(2Cl-Cbz) = 2-chlorobenzylloxycarbonyl-protected
; OTHER INFORMATION: lysine
US-09-242-724-27

Query Match 49.1%; Score 27; DB 4; Length 6;
Best Local Similarity 83.3%; Pred. No. 1.7e+05;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFF 8
: | | | | |
DB 1 EKLVEF 6

RESULT 31

US-09-242-724-30
; Sequence 30, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 30
; LENGTH: 6
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: synthetic
; OTHER INFORMATION: polypeptide
US-09-242-724-30

Query Match 49.1%; Score 27; DB 4; Length 6;
Best Local Similarity 83.3%; Pred. No. 1.7e+05;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLVFF 8
: | | | | |
DB 1 EKLVEF 6

RESULT 32

US-08-717-551A-1
; Sequence 1, Application US/08717551A
; Patent No. 6071493
; GENERAL INFORMATION:
; APPLICANT: Dana Giullian
; TITLE OF INVENTION: Identification of Agents that Protect
; TITLE OF INVENTION: Against Inflammatory Injury to Neurons
; NUMBER OF SEQUENCES: 2

; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Woodcock Washburn Kurtz Mackiewicz
; ADDRESSEE: & No. 6071493ris LLP
; STREET: One Liberty Place - 46th Floor
; CITY: Philadelphia
; STATE: PA
; COUNTRY: USA
; ZIP: 19103
; COMPUTER READABLE FORM:
; MEDIUM TYPE: DISKETTE, 3.5 INCH, 1.44 Mb STORAGE
; COMPUTER: IBM PS/2
; OPERATING SYSTEM: PC-DOS
; SOFTWARE: WORDPERFECT for WINDOWS 6.0
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/717,551A
; FILING DATE: Sept-20-96
; CLASSIFICATION: 435
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER:
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: Lori Y. Beardsell
; REGISTRATION NUMBER: 34,293
; REFERENCE/DOCKET NUMBER: BYLR-0031
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (215) 568-3100
; TELEFAX: (215) 568-3439
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 4 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; US-08-717-551A-1

Query Match 47.3%; Score 26; DB 3; Length 4;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 4; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQK 4

Db 1 HHQK 4

RESULT 33
US-09-242-724-33
; Sequence 33, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 33
; LENGTH: 6
; TYPE: PPT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: synthetic
; OTHER INFORMATION: polypeptide
US-09-242-724-33

Query Match 47.3%; Score 26; DB 4; Length 6;
Best Local Similarity 83.3%; Pred. No. 1.7e+05;
Matches 5; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QKLVFF 8

Db 1 KKLVFF 6

RESULT 34
US-08-127-904-15
; Sequence 15, Application US/08127904
; Patent No. 5470951
; GENERAL INFORMATION:
; APPLICANT: Eugene Roberts
; TITLE OF INVENTION: Method For Antagonizing
; TITLE OF INVENTION: Amnestic Effects of Amyloid n
; TITLE OF INVENTION: Protein and Improving the
; TITLE OF INVENTION: Quality of Life in Individuals
; TITLE OF INVENTION: With Alzheimer Disease
; NUMBER OF SEQUENCES: 15
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: City of Hope
; STREET: 1500 East Duarte Road
; CITY: Duarte
; STATE: California
; COUNTRY: United States of America
; ZIP: 91010-0269
; COMPUTER READABLE FORM:
; MEDIUM TYPE: 3M Double Density 5 1/4" diskette
; COMPUTER: Wang PC
; OPERATING SYSTEM: MS DOS Version 3.20
; SOFTWARE: Microsoft
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/127,904
; FILING DATE: 29 September 1993
; CLASSIFICATION: 424
; PRIOR APPLICATION DATA: NO. 5470951e
; ATTORNEY/AGENT INFORMATION:
; NAME: Irons, Edward S.
; REGISTRATION NUMBER: 16,541
; REFERENCE/DOCKET NUMBER: NO. 5470951e
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 783-6040
; TELEFAX: (202) 783-6031
; TELEX: No. 5470951e
; INFORMATION FOR SEQ ID NO: 15:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5
; TYPE: Amino Acid
; STRANDEDNESS:
; TOPOLOGY: Unknown
US-08-127-904-15
Query Match 45.5%; Score 25; DB 1; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
Db 1 KLVFF 5

RESULT 35
US-08-612-785B-10
; Sequence 10, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
; TITLE OF INVENTION: Aggregation
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:

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; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 10:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; US-08-612-785B-10

Query Match 45.5%; Score 25; DB 2; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
Db 1 KLVFF 5

RESULT 36
US-08-970-833-2
; Sequence 2, Application US/08970833
; Patent No. 6022859
; GENERAL INFORMATION:
; APPLICANT: Kiessling, Laura L.
; APPLICANT: Murphy, Regina M.
; TITLE OF INVENTION: INHIBITORS OF BETA-AMYLOID TOXICITY
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Quarles & Brady
; STREET: 411 East Wisconsin Avenue
; CITY: Milwaukee
; STATE: Wisconsin
; COUNTRY: U.S.A.
; ZIP: 53202-4497
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/970,833
; FILING DATE:
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Baker, Jean C.
; REGISTRATION NUMBER: 35,433
; REFERENCE/DOCKET NUMBER: 960296.94291
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (414) 277-5709
; TELEFAX: (414) 271-3552

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; INFORMATION FOR SEQ ID NO: 2:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; US-08-970-833-2

Query Match 45.5%; Score 25; DB 3; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
Db 1 KLVFF 5

RESULT 37
US-08-703-675C-46
; Sequence 46, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Findels, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/703,675C
; FILING DATE: 27-AUG-1996
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/616,081
; FILING DATE: 14-MAR-1996
; ATTORNEY/AGENT INFORMATION:
; NAME: Kara, Catherine J.
; REGISTRATION NUMBER: 41,106
; REFERENCE/DOCKET NUMBER: PPI-016CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 46:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; US-08-703-675C-46

Query Match 45.5%; Score 25; DB 4; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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QY 4 KLVFF 8
| | | | |
Db 1 KLVFF 5

RESULT 38

US-09-242-724-25
; Sequence 25, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 25
; LENGTH: 5
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: :
US-09-242-724-25

Query Match 45.5%; Score 25; DB 4; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
| | | | |
Db 1 KLVFF 5

RESULT 39

US-09-242-724-26
; Sequence 26, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 26
; LENGTH: 5
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; NAME/KEY: MOD_RES
; LOCATION: (1)
; OTHER INFORMATION: ACETYLATION; K(2cl-Cbz) -
; OTHER INFORMATION: 2-chlorobenzoyloxycarbonyl-protected lysine
US-09-242-724-26

Query Match 45.5%; Score 25; DB 4; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
| | | | |
Db 1 KLVFF 5

RESULT 40

US-08-617-267C-10
; Sequence 10, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:

; APPLICANT: Findels, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: Decontti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 10:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-617-267C-10

Query Match 45.5%; Score 25; DB 4; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
| | | | |
Db 1 KLVFF 5

RESULT 41

US-09-264-709A-28
; Sequence 28, Application US/09264709A
; Patent No. 6320024
; GENERAL INFORMATION:
; APPLICANT: Roberts, Eugene
; TITLE OF INVENTION: Method for Design of Substances that Enhance Memory and Improve the Quality of Life
; FILE REFERENCE: 2124-310
; CURRENT APPLICATION NUMBER: US/09/264,709A
; CURRENT FILING DATE: 1999-03-09
; PRIOR APPLICATION NUMBER: 08/797,782
; PRIOR FILING DATE: 1997-02-07
; NUMBER OF SEQ ID NOS: 39
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 28
; LENGTH: 5
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence:
; OTHER INFORMATION: memory-modulating peptide

US-09-264-709A-28

Query Match 45.5%; Score 25; DB 4; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 6 VFFAE 10
 |||||
Db 1 VFFAE 5

RESULT 42

PCT-US94-10475-15
Sequence 15, Application PC/TUS9410475

GENERAL INFORMATION:
APPLICANT: Eugene Roberts
TITLE OF INVENTION: Method For
Antagonizing Amnestic
EFFECTS OF Amyloid n
TITLE OF INVENTION: Effects of Amyloid n
TITLE OF INVENTION: Protein and Improving
the Quality of Life
TITLE OF INVENTION: in Individuals
TITLE OF INVENTION: With Alzheimer Disease
NUMBER OF SEQUENCES: 15
CORRESPONDENCE ADDRESS:
ADDRESSEE: City of Hope
STREET: 1500 East Duarte Road
CITY: Duarte
STATE: California
COUNTRY: United States of America
ZIP: 91010-0269
COMPUTER READABLE FORM:
MEDIUM TYPE: 3M Double Density 5 1/4"
MEDIUM TYPE: diskette
COMPUTER: Wang PC
OPERATING SYSTEM: MS DOS Version 3.20
SOFTWARE: Microsoft
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US94/10475
FILING DATE: 16 September 1994

CLASSIFICATION:
PRIOR APPLICATION DATA: U. S. Application
Serial No.
PRIOR APPLICATION DATA: 08/127,904; filed
PRIOR APPLICATION DATA: 29 September 1993
ATTORNEY/AGENT INFORMATION:

NAME: Irons, Edward S.
REGISTRATION NUMBER: 16,541
REFERENCE/DOCKET NUMBER: None
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202) 626-3564 or 783-6030
TELEFAX: (202) 783-6031
TELEX: None

INFORMATION FOR SEQ ID NO: 15:

SEQUENCE CHARACTERISTICS:
LENGTH: 5
TYPE: Amino Acid
STRANDEDNESS:
TOPOLOGY: Unknown

PCT-US94-10475-15

Query Match 45.5%; Score 25; DB 5; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
 |||||
Db 1 KLVFF 5

RESULT 43

US-08-612-785B-31

Sequence 31, Application US/08612785B

Patent No. 5854204

GENERAL INFORMATION:
APPLICANT: Findeis, Mark A. et al.
TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
Aggregation
NUMBER OF SEQUENCES: 40
CORRESPONDENCE ADDRESS:

ADDRESSEE: LAHIVE & COCKFIELD
STREET: 28 State Street, Suite 510
CITY: Boston
STATE: Massachusetts
COUNTRY: USA
ZIP: 02109-1875

COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk

OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/612,785B
FILING DATE: Herewith
CLASSIFICATION: 514

PRIOR APPLICATION DATA: USN 08/404,831
APPLICATION NUMBER: USN 08/404,831
FILING DATE: 14-MAR-1995

PRIOR APPLICATION DATA:

APPLICATION NUMBER: USN 08/475,579
FILING DATE: 07-JUN-1995

PRIOR APPLICATION DATA: USN 08/548,998
APPLICATION NUMBER: USN 08/548,998
FILING DATE: 27-OCT-1995

ATTORNEY/AGENT INFORMATION:

NAME: DeConti, Giulio A.
REGISTRATION NUMBER: 31,503

REFERENCE/DOCKET NUMBER: PPI-002CP3
TELECOMMUNICATION INFORMATION:

TELEPHONE: (617)227-7400

TELEFAX: (617)742-4214

INFORMATION FOR SEQ ID NO: 31:

SEQUENCE CHARACTERISTICS:

LENGTH: 6 amino acids

TYPE: amino acid

TOPOLOGY: linear

MOLECULE TYPE: peptide

FEATURE:

NAME/KEY: Modified site

LOCATION: 6

OTHER INFORMATION: /note= Xaa is beta-alanyl

US-08-612-785B-31

Query Match 45.5%; Score 25; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
 |||||
Db 1 KLVFF 5

RESULT 44

US-08-664-379B-19

Sequence 19, Application US/08664379B
Patent No. 6034211

GENERAL INFORMATION:

APPLICANT: Kelly, Jeffery W.
TITLE OF INVENTION: BETA-SHEET NUCLEATING PEPTIDOMIMETICS

NUMBER OF SEQUENCES: 19

CORRESPONDENCE ADDRESS:

ADDRESSEE: Fish & Richardson P.C.

STREET: 225 Franklin Street

CITY: Boston

STATE: MA

COUNTRY: U.S.A.

```
; ZIP: 02110-2804
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette
; COMPUTER: IBM Compatible
; OPERATING SYSTEM: DOS
; SOFTWARE: FASTSEQ Version 2.0
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/664,379B
; FILING DATE: 14-JUN-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 60/018,925
; FILING DATE: 03-JUN-1996
; ATTORNEY/AGENT INFORMATION:
; NAME: Freeman, John W.
; REGISTRATION NUMBER: 29,066
; REFERENCE/DOCKET NUMBER: 08435/003001
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617-542-5070
; TELEFAX: 617-542-8906
; TELEX: 200154
; INFORMATION FOR SEQ ID NO: 19:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; FEATURE:
; LOCATION: 1...1
; OTHER INFORMATION: wherein Xaa at position 1 is Ornithine
;
; US-08-664-379B-19

Query Match 45.5%; Score 25; DB 3; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
Db 2 KLVFF 6

RESULT 45
US-08-703-675C-44
; Sequence 44, Application US/08703675C
; Patent No. 6303567
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of -Amyloid Peptide
; NUMBER OF SEQUENCES: 46
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/703,675C
; FILING DATE: 27-AUG-1996
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; Aggregation Comprising D-
```

```
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/616,081
; FILING DATE: 14-MAR-1996
; ATTORNEY/AGENT INFORMATION:
; NAME: Kara, Catherine J.
; REGISTRATION NUMBER: 41,106
; REFERENCE/DOCKET NUMBER: PPI-016CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 44:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; FEATURE:
; NAME/KEY: Modified site
; LOCATION: 6
; OTHER INFORMATION: /note= Xaa is beta-alanyl
;
; US-08-703-675C-44

Query Match 45.5%; Score 25; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
Db 1 KLVFF 5

RESULT 46
US-09-242-724-31
; Sequence 31, Application US/09242724
; Patent No. 6316405
; GENERAL INFORMATION:
; APPLICANT: Solomon, Michael E.
; APPLICANT: Rich, Daniel H.
; TITLE OF INVENTION: Cyclosporin A Conjugates and Uses Therefor
; FILE REFERENCE: Cyclosporin Analogs
; CURRENT APPLICATION NUMBER: US/09/242,724
; CURRENT FILING DATE: 1999-02-22
; NUMBER OF SEQ ID NOS: 33
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 31
; LENGTH: 6
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: synthetic
; OTHER INFORMATION: polypeptide
;
; US-09-242-724-31

Query Match 45.5%; Score 25; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 4 KLVFF 8
Db 1 KLVFF 5

RESULT 47
US-08-617-267C-31
; Sequence 31, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
```

```

; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent In Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 31:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; FEATURE:
; NAME/KEY: Modified site
; LOCATION: 6
; OTHER INFORMATION: /note= Xaa is beta-alanyl
; US-08-617-267C-31

Query Match 45.5%; Score 25; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
Db 1 KLVFF 5

RESULT 48
US-08-617-267C-43
; Sequence 43, Application US/08617267C
; Patent No. 6319498
; GENERAL INFORMATION:
; APPLICANT: Fintel, Mark A. et al.
; TITLE OF INVENTION: Modulators of Amyloid Aggregation
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD, LLP
; STREET: 28 State Street
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent In Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/617,267C
; FILING DATE: 14-MAR-1996
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)227-5941
; INFORMATION FOR SEQ ID NO: 31:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 6 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; FEATURE:
; NAME/KEY: Modified site
; LOCATION: 6
; OTHER INFORMATION: /note= Xaa is beta-alanyl
; US-08-617-267C-31

Query Match 45.5%; Score 25; DB 4; Length 6;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
Db 1 KLVFF 5

RESULT 49
US-08-970-833-4
; Sequence 4, Application US/08970833
; Patent No. 6022859
; GENERAL INFORMATION:
; APPLICANT: Kiessling, Laura L.
; APPLICANT: Murphy, Regina M.
; TITLE OF INVENTION: INHIBITORS OF BETA-AMYLOID TOXICITY
; NUMBER OF SEQUENCES: 11
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Quarles & Brady
; STREET: 411 East Wisconsin Avenue
; CITY: Milwaukee
; STATE: Wisconsin
; COUNTRY: U.S.A.
; ZIP: 53202-4497
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent In Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/970,833
; FILING DATE:
; CLASSIFICATION: 530
; ATTORNEY/AGENT INFORMATION:
; NAME: Baker, Jean C.
; REGISTRATION NUMBER: 35,433
; REFERENCE/DOCKET NUMBER: 960296.94291
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (414) 277-5709
; TELEFAX: (414) 271-3552
; INFORMATION FOR SEQ ID NO: 4:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 8 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
; US-08-970-833-4
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Query Match 45.5%; Score 25; DB 3; Length 8;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 6 VFFAE 10
Db 1 VFFAE 5

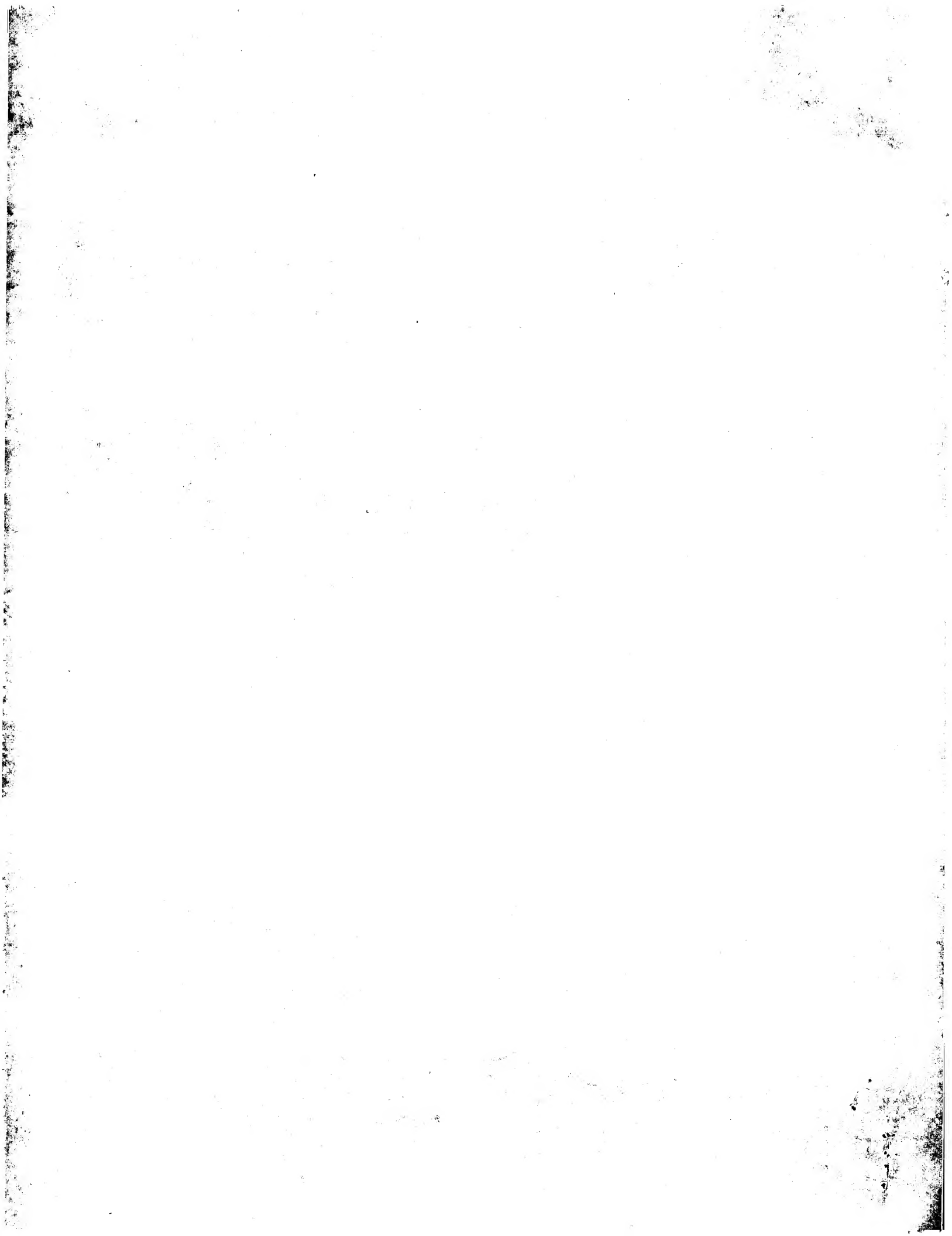
RESULT 50

US-08-612-785B-11
; Sequence 11, Application US/08612785B
; Patent No. 5854204
; GENERAL INFORMATION:
; APPLICANT: Findeis, Mark A. et al.
; TITLE OF INVENTION: Ab Peptides that Modulate b-Amyloid
; TITLE OF INVENTION: Aggregation
; NUMBER OF SEQUENCES: 40
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: LAHIVE & COCKFIELD
; STREET: 28 State Street, Suite 510
; CITY: Boston
; STATE: Massachusetts
; COUNTRY: USA
; ZIP: 02109-1875
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/612,785B
; FILING DATE: Herewith
; CLASSIFICATION: 514
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/404,831
; FILING DATE: 14-MAR-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/475,579
; FILING DATE: 07-JUN-1995
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: USSN 08/548,998
; FILING DATE: 27-OCT-1995
; ATTORNEY/AGENT INFORMATION:
; NAME: DeConti, Giulio A.
; REGISTRATION NUMBER: 31,503
; REFERENCE/DOCKET NUMBER: PPI-002CP3
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (617)227-7400
; TELEFAX: (617)742-4214
; INFORMATION FOR SEQ ID NO: 11:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 5 amino acids
; TYPE: amino acid
; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-08-612-785B-11

Query Match 43.6%; Score 24; DB 2; Length 5;
Best Local Similarity 100.0%; Pred. No. 1.7e+05;
Matches 5; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 5 LVFFA 9
Db 1 LVFFA 5

Search completed: October 29, 2002, 09:39:08
Job time : 14 secs



GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:35:42 ; Search time 14 Seconds
(without alignments)
68.635 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 283138 seqs, 96089334 residues

Total number of hits satisfying chosen parameters: 1099

Minimum DB seq length: 0
Maximum DB seq length: 10

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : PIR_71.*
1: pir1.*
2: pir2.*
3: pir3.*
4: pir4.*

Pred. No. is the number of results predicted by chance to have a
score greater than or equal to the score of the result being printed,
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	20	36.4	5	2 C41225	copper resistance
2	20	36.4	9	2 PT0080	60K Ca binding pro
3	18	32.7	10	2 S65387	cytochrome-c oxida
4	16	29.1	6	2 A46474	Fc epsilon Riib -
5	16	29.1	9	2 S13889	phosphoenolpyruvat
6	16	29.1	10	2 S74176	gluconokinase (EC
7	15	27.3	8	2 S21273	cellulase (EC 3.2.
8	15	27.3	9	2 A61102	parathyroid hormon
9	15	27.3	9	2 T31612	hypothetical prote
10	15	27.3	9	2 S10920	venom protein HR-3
11	15	27.3	9	2 B39504	octamer-binding pr
12	15	27.3	10	2 PT0310	Ig heavy chain CRD
13	15	27.3	10	2 PH0807	T-cell receptor al
14	14	25.5	7	2 A30812	sex pheromone cCF1
15	14	25.5	9	2 B20569	serum amyloid P-co
16	14	25.5	10	2 PH0113	alpha-amylyase (EC
17	14	25.5	10	2 S43631	cytochrome-c oxida
18	13	23.6	4	2 T46627	hypothetical prote
19	13	23.6	6	2 S71349	beta-crystallin B2
20	13	23.6	8	2 PT0368	Ig gamma chain C r
21	13	23.6	9	2 S55696	phosphoenolpyruvat
22	13	23.6	10	2 S65388	cytochrome-c oxida
23	13	23.6	10	2 S30348	clotting protein -
24	13	23.6	10	2 S43625	cytochrome-c oxida
25	13	23.6	10	2 PT0284	Ig heavy chain CRD
26	13	23.6	10	2 B45482	platelet activatin
27	13	23.6	10	2 T13838	cytochrome-c oxida
28	13	23.6	10	2 T13976	cytochrome-c oxida
29	13	23.6	10	2 T17057	cytochrome-c oxida

30	13	23.6	10	2 T12303	cytochrome-c oxida
31	13	23.6	10	2 T14019	cytochrome-c oxida
32	13	23.6	10	2 T17060	cytochrome-c oxida
33	13	23.6	10	2 T14043	cytochrome-c oxida
34	13	23.6	10	2 T14054	cytochrome-c oxida
35	13	23.6	10	2 T17066	cytochrome-c oxida
36	13	23.6	10	2 T17069	cytochrome-c oxida
37	13	23.6	10	2 T12308	cytochrome-c oxida
38	13	23.6	10	2 T17072	cytochrome-c oxida
39	13	23.6	10	2 T12312	cytochrome-c oxida
40	13	23.6	10	2 T12316	cytochrome-c oxida
41	13	23.6	10	2 T12321	cytochrome-c oxida
42	13	23.6	10	2 T14219	cytochrome-c oxida
43	12	21.8	4	2 J01273	neuropeptide Antho
44	12	21.8	4	2 A32480	achatin-1 - giant
45	12	21.8	6	2 A60986	N-formyl oligopept
46	12	21.8	6	2 I59142	platelet-derived g
47	12	21.8	6	2 A43129	neuropeptide GNFFR
48	12	21.8	7	2 PT0246	Ig heavy chain CRD
49	12	21.8	7	2 I48668	alpha-myosin heavy
50	12	21.8	8	2 T13818	cytochrome oxidase

ALIGNMENTS

RESULT 1

C41225
copper resistance protein - Pseudomonas syringae pv. tomato (fragment)
C;Species: Pseudomonas syringae pv. tomato
C;Date: 19-Jun-1992 #sequence_revision 19-Jun-1992 #text_change 24-Jun-1993
C;Accession: C41225
R;Cha, J.S.; Cooksey, D.A.
Proc. Natl. Acad. Sci. U.S.A. 88, 8915-8919, 1991
A;Title: Copper resistance in Pseudomonas syringae mediated by periplasmic and outer
A;Reference number: A41225; MUID:92020961
A;Accession: C41225
A;Status: preliminary
A;Molecule type: protein
A;Residues: 1-5 <CHA>

Query Match 36.4%; Score 20; DB 2; Length 5;
Best Local Similarity 80.0%; Pred. No. 2.8e+05;
Matches 4; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKLV 6
| | | |
Db 1 HPKLV 5

RESULT 2

PT0080
60K Ca binding protein - edible frog (fragment)
C;Species: Rana esculenta (edible frog)
C;Date: 16-Jul-1999 #sequence_revision 16-Jul-1999 #text_change 16-Jul-1999
C;Accession: PT0080
R;Treveso, S.; Zorzato, F.; Chiozzi, P.; Melandri, P.; Volpe, P.; Pozzan, T.
Biochem. Biophys. Res. Commun. 175, 444-450, 1991
A;Title: Frog brain expresses a 60 kDa Ca2+ binding protein similar to mammalian calr
A;Reference number: PT0080; MUID:91207333
A;Accession: PT0080
A;Status: preliminary
A;Molecule type: protein
A;Residues: 1-9 <TRE>

Query Match 36.4%; Score 20; DB 2; Length 9;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 4; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8
| | | |
Db 3 LVFF 6

RESULT 3

S65387
 cytochrome-c oxidase (EC 1.9.3.1) chain VII b, cardiac - rat (fragment)
 C:Species: Rattus norvegicus (Norway rat)
 C:Date: 12-Feb-1998 #sequence_revision 20-Feb-1998 #text_change 16-Jul-1999
 C:Accession: S65387; S65386
 R:Schaeffer, H.; Noack, H.; Halangk, W.; Brandt, U.; von Jagow, G.
 Eur. J. Biochem. 230, 235-241, 1995
 A:Title: Cytochrome-c oxidase in developing rat heart. Enzymic properties and amino-term
 A:Reference number: S65372; MUID:95324529
 A:Accession: S65387
 A:Status: preliminary
 A:Molecule type: protein
 A:Residues: 1-10 <SCH>
 A:Accession: S65386
 A:Status: preliminary
 A:Molecule type: protein
 A:Residues: 1-10 <SC2>
 C:Keywords: cardiac muscle; heart; oxidoreductase

Query Match 32.7%; Score 18; DB 2; Length 10;
 Best Local Similarity 100.0%; Pred. No. 1.7e+03;
 Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQK 4
 |||
 Db 2 HQK 4

RESULT 4

A46474
 Fc epsilon RIIB - mouse (fragment)
 C:Species: Mus musculus (house mouse)
 C:Date: 18-Jun-1993 #sequence_revision 18-Nov-1994 #text_change 11-Apr-1995
 C:Accession: A46474
 R:Richards, M.L.; Katz, D.H.; Liu, F.T.
 J. Immunol. 147, 1067-1074, 1991
 A:Title: Complete genomic sequence of the murine low affinity Fc receptor for IgE. Demon
 A:Reference number: A46474; MUID:91318149
 A:Accession: A46474
 A:Status: preliminary; not compared with conceptual translation
 A:Molecule type: nucleic acid
 A:Residues: 1-6 <RIC>
 A:Experimental source: BALB C, splenic B cells
 A:Note: sequence extracted from NCBI backbone (NCBIP:45428)

Query Match 29.1%; Score 16; DB 2; Length 6;
 Best Local Similarity 100.0%; Pred. No. 2.8e+05;
 Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HH 2
 |||
 Db 4 HH 5

RESULT 5

S13889
 phosphoenolpyruvate carboxylase (EC 4.1.1.31) - maize
 C:Species: Zea mays (maize)
 C:Date: 19-Mar-1997 #sequence_revision 19-Mar-1997 #text_change 01-Aug-1997
 C:Accession: S13889
 R:Jiao, J.; Chollet, R.
 Arch. Biochem. Biophys. 283, 300-305, 1990
 A:Title: Regulatory phosphorylation of serine-15 in maize phosphoenolpyruvate carboxylas
 A:Reference number: S13889; MUID:91112741
 A:Accession: S13889
 A:Status: preliminary
 A:Molecule type: protein
 A:Residues: 1-9 <JIA>
 C:Keywords: carbon-carbon lyase; carboxy-lyase

Query Match 29.1%; Score 16; DB 2; Length 9;

Best Local Similarity 100.0%; Pred. No. 2.8e+05;
 Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HH 2
 |||
 Db 1 HH 2

RESULT 6

S74176
 gluconokinase (EC 2.7.1.12), thermoresistant - Escherichia coli (fragment)
 C:Species: Escherichia coli
 C:Date: 14-Apr-1998 #sequence_revision 24-Apr-1998 #text_change 07-May-1999
 C:Accession: S74176
 R:Izu, H.; Adachi, O.; Yamada, M.
 FEBS Lett. 394, 14-16, 1996
 A:Title: Purification and characterization of the Escherichia coli thermoresistant gl
 A:Reference number: S74176; MUID:97074194
 A:Accession: S74176
 A:Molecule type: protein
 A:Residues: 1-10 <IZU>
 A:Experimental source: strain K-12
 C:Genetics:
 A:Gene: gntK
 C:Keywords: dimer; phosphotransferase

Query Match 29.1%; Score 16; DB 2; Length 10;
 Best Local Similarity 100.0%; Pred. No. 4.3e+03;
 Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HH 2
 |||
 Db 7 HH 8

RESULT 7

S21273
 cellulase (EC 3.2.1.4) - Clostridium thermocellum (fragment)
 N:Alternate names: endo-1,4-beta-glucanase
 C:Species: Clostridium thermocellum
 C:Date: 04-Dec-1992 #sequence_revision 04-Dec-1992 #text_change 22-Nov-1996
 C:Accession: S21273
 R:Romanec, M.P.M.; Fauth, U.; Kobayashi, T.; Huskisson, N.S.; Barker, P.J.; Demain,
 Biochem. J. 283, 69-73, 1992
 A:Title: Purification and characterization of a new endoglucanase from Clostridium th
 A:Reference number: S21273; MUID:92231850
 A:Accession: S21273
 A:Molecule type: protein
 A:Residues: 1-8 <ROM>
 C:Function:
 A:Description: hydrolysis of 1,4-beta-D-glucosidic linkages in beta-D-glucans such as
 A:Pathway: cellulose degradation
 C:Keywords: glycosidase; hydrolase; polysaccharide degradation

Query Match 27.3%; Score 15; DB 2; Length 8;
 Best Local Similarity 100.0%; Pred. No. 2.8e+05;
 Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
 |||
 Db 4 FAE 6

RESULT 8

A61102
 parathyroid hormone-like protein, humoral hypercalcemia of malignancy - dog (fragment)
 C:Species: Canis lupus familiaris (dog)
 C:Date: 31-Dec-1993 #sequence_revision 31-Dec-1993 #text_change 17-Mar-1999
 C:Accession: A61102
 R:Weir, E.C.; Burtis, W.J.; Morris, C.A.; Brady, T.G.; Insogna, K.L.
 Endocrinology 123, 2744-2751, 1988
 A:Title: Isolation of 16,000-dalton parathyroid hormone-like proteins from two animal
 A:Reference number: A61102; MUID:89064600

A:Accession: A61102
A:Molecule type: protein

A:Residues: 1-9 <WEI>

A:Experimental source: apocrine cell adenocarcinoma

C:Superfamily: parathyroid hormone-related protein; parathyroid hormone homology

C:Keywords: hormone; humoral hypercalcemia

Query Match 27.3%; Score 15; DB 2; Length 9;
Best Local Similarity 75.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKL 5

|||

Db 6 HQLL 9

RESULT 9

T31612

hypothetical protein Y50E8A.h - Caenorhabditis elegans

C:Species: Caenorhabditis elegans

C:Date: 29-Oct-1999 #sequence_revision 29-Oct-1999 #text_change 29-Oct-1999

C:Accession: T31612

R:Steward, C.

submitted to the EMBL Data Library, September 1999

A:Reference number: Z21047

A:Accession: T31612

A>Status: preliminary; translated from GB/EMBL/DDBJ

A:Molecule type: DNA

A:Residues: 1-9 <WIL>

A:Cross-references: EMBL:AL117200; NID:el549770; PIDN:CAB55051.1; CESP:Y50E8A.h

A:Experimental source: clone Y50E8A

C:Genetics:

A:Gene: CBSP:Y50E8A.h

Query Match 27.3%; Score 15; DB 2; Length 9;
Best Local Similarity 50.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 1 HHOK 4

|||

Db 5 HREK 8

RESULT 10

S10920

venom protein HR-3 - oriental hornet (fragment)

C:Species: Vespa orientalis (oriental hornet)

C:Date: 29-Jan-1993 #sequence_revision 29-Jan-1993 #text_change 08-Dec-1995

C:Accession: S10920

R:Tsuchibae, M.U.; Akhmedova, N.U.; Kazakov, I.; Korneev, A.S.; Gagel'gans, A.I.

Biochemistry (N.Y.) 53, 183-190, 1988

A:Title: Low-molecular-weight peptides of venom of the giant hornet Vespa orientalis. St

A:Reference number: S06445

A:Accession: S10920

A:Molecule type: protein

A:Residues: 1-9 <TUI>

C:Keywords: venom

Query Match 27.3%; Score 15; DB 2; Length 9;
Best Local Similarity 60.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HOKLV 6

|||

Db 4 HEFLV 8

RESULT 11

B39504

octamer-binding protein, Ku-like, 83k chain - human (fragment)

C:Species: Homo sapiens (man)

C:Date: 30-Dec-1991 #sequence_revision 30-Dec-1991 #text_change 30-Sep-1993

C:Accession: B39504

R:May, G.; Sutton, C.; Gould, H.
J. Biol. Chem. 266, 3052-3059, 1991

A:Title: Purification and characterization of Ku-2, an octamer-binding protein relate

A:Reference number: A39504; MUID:91131605

A:Accession: B39504

A>Status: preliminary

A:Molecule type: protein

A:Residues: 1-9 <MAY>

Query Match 27.3%; Score 15; DB 2; Length 9;
Best Local Similarity 50.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 5 LVFFAE 10

|||

Db 1 MVFMNE 6

RESULT 12

PT0310

Ig heavy chain CRD3 region (clone 6-97) - human (fragment)

C:Species: Homo sapiens (man)

C:Date: 30-Sep-1993 #sequence_revision 30-Sep-1993 #text_change 16-Aug-1996

C:Accession: PT0310

R:Yamada, M.; Wasserman, R.; Reichard, B.A.; Shane, S.; Caton, A.J.; Rovera, G.

J. Exp. Med. 173, 395-407, 1991

A:Title: Preferential utilization of specific immunoglobulin heavy chain diversity an

A:Reference number: PT0222; MUID:91108337

A:Accession: PT0310

A:Molecule type: DNA

A:Residues: 1-10 <YAM>

A:Experimental source: B lymphocyte

C:Keywords: heterotetramer; immunoglobulin

Query Match 27.3%; Score 15; DB 2; Length 10;
Best Local Similarity 75.0%; Pred. No. 6.8e+03;
Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8

|||

Db 3 LVWF 6

RESULT 13

PH0807

T-cell receptor alpha chain (J4) - mouse (fragment)

C:Species: Mus musculus (house mouse)

C:Date: 17-Jul-1992 #sequence_revision 17-Jul-1992 #text_change 30-May-1997

C:Accession: PH0807

R:Casanova, J.L.; Romero, P.; Widmann, C.; Kourilsky, P.; Maryanski, J.L.

J. Exp. Med. 174, 1371-1383, 1991

A:Title: T cell receptor genes in a series of class I major histocompatibility comple

allelic exclusion and antigen-specific repertoire.

A:Reference number: PH0746; MUID:92078846

A:Accession: PH0807

A:Molecule type: mRNA

A:Residues: 1-10 <CAS>

A:Cross-references: EMBL:X60916

A:Experimental source: T lymphocyte

C:Keywords: T-cell receptor

Query Match 27.3%; Score 15; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 6.8e+03;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 6 VFF 8

|||

Db 7 IFF 9

RESULT 14

A30812

sex pheromone cCF10 - Enterococcus faecalis

```

C:Species: Enterococcus faecalis
C:Date: 28-Feb-1990 #sequence_revision 28-Feb-1990 #text_change 18-Jun-1993
C:Accession: A30812
R:Mori, M.; Sakagami, Y.; Ishii, Y.; Isogai, A.; Kitada, C.; Fujino, M.; Adsit, J.C.; Du
J. Biol. Chem. 263, 14574-14578, 1988
A:Title: Structure of cCf10, a peptide sex pheromone which induces conjugative transfer
A:Reference number: A30812; MUID:89008313
A:Accession: A30812
A:Molecule type: protein
A:Residues: 1-7 <MOR>

Query Match      25.5%; Score 14; DB 2; Length 7;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVF 7
Db 4 LVF 6

RESULT 15
B20569
serum amyloid P-component - smooth dogfish (fragment)
C:Species: Mustelus canis (smooth dogfish)
C:Date: 05-Jun-1987 #sequence_revision 05-Jun-1987 #text_change 18-Jun-1993
C:Accession: B20569; A05074
R:Robey, F.A.; Tanaka, T.; Liu, T.Y.
J. Biol. Chem. 258, 3889-3894, 1983
A:Title: Isolation and characterization of two major serum proteins from the dogfish, M
A:Reference number: A92419; MUID:83160932
A:Accession: B20569
A:Molecule type: protein
A:Residues: 1-9 <ROB>
C:Keywords: amyloid

Query Match      25.5%; Score 14; DB 2; Length 9;
Best Local Similarity 40.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 3 QKLVF 7
Db 5 KSLIF 9

RESULT 16
PH0113
alpha-amylase (EC 3.2.1.1) III - rice (fragment)
C:Species: Oryza sativa (rice)
C:Date: 17-Apr-1993 #sequence_revision 17-Apr-1993 #text_change 07-May-1999
C:Accession: PH0113
R:Chiba, Y.; Niede, Y.; Nakajima, T.; Ichishima, E.
Agric. Biol. Chem. 55, 901-902, 1991
A:Title: Unique enzymatic properties of alpha-amylase-III from suspension-cultured rice
A:Reference number: PH0113; MUID:91234351
A:Accession: PH0113
A:Molecule type: protein
A:Residues: 1-10 <CHI>
A:Experimental source: cv. Sasanishiki
C:Function:
A:Description: catalyzes the hydrolysis of internal 1,4-alpha-D-glucosidic bonds
A:Pathway: glycogen/starch degradation
C:Keywords: glycosidase; hydrolase; polysaccharide degradation

Query Match      25.5%; Score 14; DB 2; Length 10;
Best Local Similarity 75.0%; Pred. No. 1.1e+04;
Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKLV 6
Db 6 QELV 9

RESULT 17
S43631
cytochrome-c oxidase (EC 1.9.3.1) chain VIIa, cardiac - rainbow trout (fragment)
C:Species: Oncorhynchus mykiss (rainbow trout)
C:Date: 20-Oct-1994 #sequence_revision 01-Nov-1996 #text_change 16-Jul-1999
C:Accession: S43631
R:Freund, R.; Kadenbach, B.
Eur. J. Biochem. 221, 1111-1116, 1994
A:Title: Identification of tissue-specific isoforms for subunits Vb and VIIa of cyto
A:Reference number: S43624; MUID:94237150
A:Accession: S43631
A:Molecule type: protein
A:Residues: 1-10 <PRE>
A:Note: the source is designated as Salmo gairdneri
C:Genetics:
A:Genome: nuclear
C:Keywords: cardiac muscle; heart; membrane-associated complex; mitochondrion; oxidor

Query Match      25.5%; Score 14; DB 2; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 OKL 5
Db 8 OKL 10

RESULT 18
T46627
hypothetical protein c4 - loblolly pine
C:Species: Pinus taeda (loblolly pine)
C:Date: 18-Feb-2000 #sequence_revision 18-Feb-2000 #text_change 18-Feb-2000
C:Accession: T46627
R:Chang, S.; Puryea, J.; Funkhouser, E.A.; Newton, R.J.; Cairney, J.
submitted to the EMBL Data Library, July 1995
A:Description: Cloning of a chitinase homolog which lacks chitin binding sites and is
A:Reference number: Z23105
A:Accession: T46627
A>Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: mRNA
A:Residues: 1-4 <CHA>
A:Cross-references: EMBL:U31309; NID:9974285; PID:9974292
A:Experimental source: strain s6PTxs6PT3; 8 month seedlings

Query Match      23.6%; Score 13; DB 2; Length 4;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLV 6
Db 2 KLV 4

RESULT 19
S71349
beta-crystallin B2 - rat (fragment)
C:Species: Rattus norvegicus (Norway rat)
C:Date: 29-Jan-1998 #sequence_revision 06-Feb-1998 #text_change 07-May-1999
C:Accession: S71349
R:Dirks, R.P.H.; Kraft, H.J.; van Genesen, S.T.; Klok, E.J.; Pfundt, R.; Schoenmakers
Eur. J. Biochem. 239, 23-32, 1996
A:Title: the cooperation between two silencers creates an enhancer element that contr
A:Reference number: S71349; MUID:96305362
A:Accession: S71349
A>Status: translation not shown
A:Molecule type: DNA
A:Residues: 1-6 <DIR>
A:Cross-references: EMBL:X83671
A:Experimental source: strain Wistar; lens epithelial cells
C:Genetics:
A:Gene: CRYBB2

Query Match      23.6%; Score 13; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;

```

Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HQ 3

||

Db 5 HQ 6

RESULT 20

PT0368

Ig gamma chain C region (gamma-1) - human (fragment)

C:Species: Homo sapiens (man)

C:Date: 31-Mar-1992 #sequence_revision 31-Mar-1992 #text_change 16-Aug-1996

C:Accession: PT0368

R:Milili, M.; Fougereau, M.; Guglielmi, P.; Schiff, C.

Mol. Immunol. 28, 753-761, 1991

A:Title: Early occurrence of immunoglobulin isotype switching in human fetal liver.

A:Reference number: PT0368; MUID:91312348

A:Accession: PT0368

A:Molecule type: mRNA

A:Residues: 1-8 <MIL>

A:Experimental source: fetal liver

C:Keywords: immunoglobulin

Query Match 23.6%; Score 13; DB 2; Length 8;

Best Local Similarity 100.0%; Pred. No. 2.8e+05;

Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HQ 3

||

Db 2 HQ 3

RESULT 21

S55696

phosphoenolpyruvate carboxykinase - Trypanosoma brucei

C:Species: Trypanosoma brucei

C:Date: 28-Oct-1995 #sequence_revision 03-Nov-1995 #text_change 07-May-1999

C:Accession: S55696

R:Hunt, M.; Koehler, P.

Biochim. Biophys. Acta 1249, 15-22, 1995

A:Title: Purification and characterization of phosphoenolpyruvate carboxykinase from Try

A:Reference number: S55696; MUID:95284106

A:Accession: S55696

A>Status: preliminary

A:Molecule type: protein

A:Residues: 1-9 <HUN>

Query Match 23.6%; Score 13; DB 2; Length 9;

Best Local Similarity 50.0%; Pred. No. 2.8e+05;

Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HOKL 5

||

Db 5 HKNL 8

RESULT 22

S65388

cytochrome-c oxidase (EC 1.9.3.1) chain VII c, hepatic - rat (fragment)

C:Species: Rattus norvegicus (Norway rat)

C:Date: 28-Oct-1996 #sequence_revision 13-Mar-1997 #text_change 07-May-1999

C:Accession: S65388; S65389

R:Schaegger, H.; Noack, H.; Halangk, W.; Brandt, U.; von Jagow, G.

Eur. J. Biochem. 230, 235-241, 1995

A:Title: Cytochrome-c oxidase in developing rat heart. Enzymic properties and amino-term

A:Reference number: S65372; MUID:95324529

A:Accession: S65388

A>Status: preliminary

A:Molecule type: protein

A:Residues: 1-10 <SCH>

A:Accession: S65389

A>Status: preliminary

A:Molecule type: protein

A:Residues: 1-10 <SC2>

C:Superfamily: cytochrome-c oxidase chain VIIC

C:Keywords: Oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;

Best Local Similarity 25.0%; Pred. No. 1.7e+04;

Matches 1; Conservative 3; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHOK 4

||:::

Db 2 HYE 5

RESULT 23

S30348

clotting protein - signal crayfish

C:Species: Pacifastacus leniusculus (signal crayfish)

C:Date: 19-Mar-1997 #sequence_revision 19-Mar-1997 #text_change 17-Mar-1999

C:Accession: S30348

R:Kopacek, P.; Hall, M.; Soederhaell, K.

Eur. J. Biochem. 213, 591-597, 1993

A:Title: Characterization of a clotting protein, isolated from plasma of the freshwater

A:Reference number: S30348; MUID:95238739

A:Accession: S30348

A>Status: preliminary

A:Molecule type: protein

A:Residues: 1-10 <KOP>

Query Match 23.6%; Score 13; DB 2; Length 10;

Best Local Similarity 33.3%; Pred. No. 1.7e+04;

Matches 2; Conservative 1; Mismatches 3; Indels 0; Gaps 0;

Qy 2 HOKLVF 7

||

Db 2 HSNLEY 7

RESULT 24

S43625

cytochrome-c oxidase (EC 1.9.3.1) chain Va, hepatic - rainbow trout (fragment)

C:Species: Oncorhynchus mykiss (rainbow trout)

C:Date: 20-Oct-1994 #sequence_revision 01-Nov-1996 #text_change 18-Jul-1997

C:Accession: S43625

R:Freund, R.; Kadenbach, B.

Eur. J. Biochem. 221, 1111-1116, 1994

A:Title: Identification of tissue-specific isoforms for subunits Vb and VIa of cyto

A:Reference number: S43624; MUID:94237150

A:Accession: S43625

A:Molecule type: protein

A:Residues: 1-10 <FRE>

A:Note: the source is designated as Salmo gairdneri

C:Genetics:

A:Genome: nuclear

C:Keywords: liver; membrane-associated complex; mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;

Best Local Similarity 50.0%; Pred. No. 1.7e+04;

Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HOKL 5

||

Db 2 HAKV 5

RESULT 25

PT0284

Ig heavy chain CRD3 region (clone 4-97) - human (fragment)

C:Species: Homo sapiens (man)

C:Date: 30-Sep-1993 #sequence_revision 30-Sep-1993 #text_change 16-Aug-1996

C:Accession: PT0284

R:Yamada, M.; Wasserman, R.; Reichard, B.A.; Shane, S.; Caton, A.J.; Rovera, G.

J. Exp. Med. 173, 395-407, 1991

A:Title: Preferential utilization of specific immunoglobulin heavy chain diversity an

A:Reference number: PT0222; MUID:91108337
 A:Accession: PT0284
 A:Molecule type: DNA
 A:Residues: 1-10 <YAM>
 C:Experimental source: B lymphocyte
 C:Keywords: heterotetramer; immunoglobulin

Query Match 23.6%; Score 13; DB 2; Length 10;
 Best Local Similarity 50.0%; Pred. No. 1.7e+04;
 Matches 3; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 3 QKLVFF 8
 |||
 Db 3 QQLANF 8

RESULT 26

B45482
 platelet activating factor acetylhydrolase - human (fragment)
 C:Species: Homo sapiens (man)
 C:Date: 05-May-1995 #sequence_revision 05-May-1995 #text_change 05-May-1995
 C:Accession: B45482
 R:Stafforini, D.M.; Rollins, E.N.; Prescott, S.M.; McIntyre, T.M.
 J. Biol. Chem. 268, 3857-3865, 1993
 A:Title: The platelet-activating factor acetylhydrolase from human erythrocytes. Purification and characterization of the cDNA
 A:Reference number: A45482; MUID:93179380
 A:Accession: B45482
 A:Status: preliminary
 A:Molecule type: protein
 A:Residues: 1-10 <STA>

Query Match 23.6%; Score 13; DB 2; Length 10;
 Best Local Similarity 75.0%; Pred. No. 1.7e+04;
 Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 5 LVVF 8
 |||
 Db 3 LVVF 6

RESULT 27

T13838
 cytochrome-c oxidase (EC 1.9.3.1) chain I - Bipes biporus mitochondrion (fragment)
 C:Species: mitochondrion Bipes biporus
 C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 11-May-2000
 C:Accession: T13838
 R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.
 Mol. Biol. Evol. 14, 91-104, 1997
 A:Title: Two novel gene orders and the role of light-strand replication in rearrangement of the mitochondrial cytochrome-c oxidase subunit I
 A:Reference number: Z17789; MUID:97153826
 A:Accession: T13838
 A:Status: preliminary; translated from GB/EMBL/DBJ
 A:Molecule type: DNA
 A:Residues: 1-10 <MAC>
 A:Cross-references: EMBL:U71335; NID:g1753232; PID:g1753235; PIDN:AAB48271.1
 C:Genetics:
 A:Genome: mitochondrion
 A:Note: COI
 C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
 Best Local Similarity 66.7%; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
 |||
 Db 7 FFS 9

RESULT 28

T13976
 cytochrome-c oxidase (EC 1.9.3.1) chain I - Cnemidophorus tigris mitochondrion (fragment)
 C:Species: mitochondrion Cnemidophorus tigris

C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 11-May-2000
 C:Accession: T13976
 R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.
 Mol. Biol. Evol. 14, 91-104, 1997
 A:Title: Two novel gene orders and the role of light-strand replication in rearrangement of the mitochondrial cytochrome-c oxidase subunit I
 A:Reference number: Z17789; MUID:97153826
 A:Accession: T13976
 A:Status: preliminary; translated from GB/EMBL/DBJ
 A:Molecule type: DNA
 A:Residues: 1-10 <MAC>
 A:Cross-references: EMBL:U71332; NID:g1753236; PID:g1753239; PIDN:AAB48274.1
 C:Genetics:
 A:Genome: mitochondrion
 A:Note: COI
 C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
 Best Local Similarity 66.7%; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
 |||
 Db 7 FFS 9

RESULT 29

T17057
 cytochrome-c oxidase (EC 1.9.3.1) chain I - Crotophytus collaris mitochondrion (fragment)
 C:Species: mitochondrion Crotophytus collaris
 C:Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 22-Oct-1999
 C:Accession: T17057
 R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Papenfuss, T.J.
 J. Mol. Evol. 44, 660-674, 1997
 A:Title: Evolutionary shifts in three major structural features of the mitochondrial cytochrome-c oxidase subunit I
 A:Reference number: Z18674; MUID:97315309
 A:Accession: T17057
 A:Status: preliminary; translated from GB/EMBL/DBJ
 A:Molecule type: DNA
 A:Residues: 1-10 <MAC>
 A:Cross-references: EMBL:U82681; NID:g3603108; PID:g3603111; PIDN:AAC62272.1
 C:Genetics:
 A:Genome: mitochondrion
 A:Note: COI
 C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
 Best Local Similarity 66.7%; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
 |||
 Db 7 FFS 9

RESULT 30

T12303
 cytochrome-c oxidase (EC 1.9.3.1) chain I - Dipsosaurus dorsalis mitochondrion (fragment)
 C:Species: mitochondrion Dipsosaurus dorsalis
 C:Date: 23-Jul-1999 #sequence_revision 23-Jul-1999 #text_change 22-Oct-1999
 C:Accession: T12303
 R:Schulte, J.A.; Macey, J.R.; Larson, A.; Papenfuss, T.J.
 Mol. Phylogenet. Evol. 10, 367-376, 1998
 A:Title: Molecular tests of phylogenetic taxonomies: A general procedure and example
 A:Reference number: Z17488; MUID:99162288
 A:Accession: T12303
 A:Status: preliminary; translated from GB/EMBL/DBJ
 A:Molecule type: DNA
 A:Residues: 1-10 <SCH>
 A:Cross-references: EMBL:AF049857; NID:g4105726; PID:g4105729; PIDN:AAD02514.1
 C:Genetics:
 A:Genome: mitochondrion
 A:Note: COI
 C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
||:
Db 7 FFS 9

RESULT 31

Tl14019

cytochrome-c oxidase (EC 1.9.3.1) chain I - Eremlas grammica mitochondrion (fragment)

C:Species: mitochondrion Eremlas grammica

C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 11-May-2000

C:Accession: Tl14019

R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.

Mol. Biol. Evol. 14, 91-104, 1997

A:Title: Two novel gene orders and the role of light-strand replication in rearrangement

A:Reference number: Z17789; MUID:97153826

A:Accession: Tl14019

A:Status: preliminary; translated from GB/EMBL/DBJ

A:Molecule type: DNA

A:Residues: 1-10 <MAC>

A:Cross-references: EMBL:U71331; NID:gl753240; PID:gl753243; PIDN:AAB48277.1

C:Genetics:

A:Genome: mitochondrion

A:Note: COI

C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
||:
Db 7 FFS 9

RESULT 32

Tl17060

cytochrome-c oxidase (EC 1.9.3.1) chain I - Gambellia wislizenii mitochondrion (fragment)

C:Species: mitochondrion Gambellia wislizenii

C:Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 22-Oct-1999

C:Accession: Tl17060

R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Papenfuss, T.J.

J. Mol. Evol. 44, 660-674, 1997

A:Title: Evolutionary shifts in three major structural features of the mitochondrial gen

A:Reference number: Z18674; MUID:97315309

A:Accession: Tl17060

A:Status: preliminary; translated from GB/EMBL/DBJ

A:Molecule type: DNA

A:Residues: 1-10 <MAC>

A:Cross-references: EMBL:U82682; NID:g3603120; PID:g3603123; PIDN:AAC62281.1

C:Genetics:

A:Genome: mitochondrion

A:Note: COI

C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
||:
Db 7 FFS 9

RESULT 33

Tl14043

cytochrome-c oxidase (EC 1.9.3.1) chain I - Lialis jicari mitochondrion (fragment)

C:Species: mitochondrion Lialis jicari

C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 11-May-2000

C:Accession: Tl14043
R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.
Mol. Biol. Evol. 14, 91-104, 1997

A:Title: Two novel gene orders and the role of light-strand replication in rearrangem

A:Reference number: Z17789; MUID:97153826

A:Accession: Tl14043

A:Status: preliminary; translated from GB/EMBL/DBJ

A:Molecule type: DNA

A:Residues: 1-10 <MAC>

A:Cross-references: EMBL:U71327; NID:gl753244; PID:gl753247; PIDN:AAB48280.1

C:Genetics:

A:Genome: mitochondrion

A:Note: COI

C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
||:
Db 7 FFS 9

RESULT 34

Tl14054

cytochrome-c oxidase (EC 1.9.3.1) chain I - Mabuya aurata mitochondrion (fragment)

C:Species: mitochondrion Mabuya aurata

C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 11-May-2000

C:Accession: Tl14054

R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.

Mol. Biol. Evol. 14, 91-104, 1997

A:Title: Two novel gene orders and the role of light-strand replication in rearrangem

A:Reference number: Z17789; MUID:97153826

A:Accession: Tl14054

A:Status: preliminary; translated from GB/EMBL/DBJ

A:Molecule type: DNA

A:Residues: 1-10 <MAC>

A:Cross-references: EMBL:U71330; NID:gl753248; PID:gl753251; PIDN:AAB48283.1

C:Genetics:

A:Genome: mitochondrion

A:Note: COI

C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
||:
Db 7 FFS 9

RESULT 35

Tl17066

cytochrome-c oxidase (EC 1.9.3.1) chain I - Oplurus cuvieri mitochondrion (fragment)

C:Species: mitochondrion Oplurus cuvieri

C:Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 22-Oct-1999

C:Accession: Tl17066

R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Papenfuss, T.J.

J. Mol. Evol. 44, 660-674, 1997

A:Title: Evolutionary shifts in three major structural features of the mitochondrial

A:Reference number: Z18674; MUID:97315309

A:Accession: Tl17066

A:Status: preliminary; translated from GB/EMBL/DBJ

A:Molecule type: DNA

A:Residues: 1-10 <MAC>

A:Cross-references: EMBL:U82685; NID:g3603136; PID:g3603139; PIDN:AAC62293.1

C:Genetics:

A:Genome: mitochondrion

A:Note: COI

C:Keywords: mitochondrion; oxidoreductase

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Query Match      23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
   ||:
Db 7 FFS 9

RESULT 36
Tl7069
cytochrome-c oxidase (EC 1.9.3.1) chain I - Phrynosoma douglassii mitochondrion (fragment)
C:Species: mitochondrion Phrynosoma douglassii
C>Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 22-Oct-1999
C:Accession: Tl7069
R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Papenfuss, T.J.
J. Mol. Evol. 44, 660-674, 1997
A:Title: Evolutionary shifts in three major structural features of the mitochondrial genome
A:Reference number: Z18674; MUID:97315309
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Accession: Tl7069
A:Molecule type: DNA
A:Residues: 1-10 <MAC>
A:Cross-references: EMBL:U82687; NID:g3603144; PID:g3603147; PIDN:AAC62299.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match      23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
   ||:
Db 7 FFS 9

RESULT 37
Tl2308
cytochrome-c oxidase (EC 1.9.3.1) chain I - Sator angustus mitochondrion (fragment)
C:Species: mitochondrion Sator angustus
C>Date: 23-Jul-1999 #sequence_revision 23-Jul-1999 #text_change 22-Oct-1999
C:Accession: Tl2308
R:Schulte, J.A.; Macey, J.R.; Larson, A.; Papenfuss, T.J.
Mol. Phylogenet. Evol. 10, 367-376, 1998
A:Title: Molecular tests of phylogenetic taxonomies: A general procedure and example
A:Reference number: Z17488; MUID:99162288
A:Accession: Tl2308
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <SCH>
A:Cross-references: EMBL:AF049859; NID:g4105734; PID:g4105737; PIDN:AAD02520.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match      23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
   ||:
Db 7 FFS 9

RESULT 38
Tl7072
cytochrome-c oxidase (EC 1.9.3.1) chain I - Sauromalus obesus mitochondrion (fragment)
C:Species: mitochondrion Sauromalus obesus
C>Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 22-Oct-1999
C:Accession: Tl7072

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R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Papenfuss, T.J.
J. Mol. Evol. 44, 660-674, 1997
A:Title: Evolutionary shifts in three major structural features of the mitochondrial genome
A:Reference number: Z18674; MUID:97315309
A:Accession: Tl7072
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <MAC>
A:Cross-references: EMBL:U82687; NID:g3603152; PID:g3603155; PIDN:AAC62305.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match      23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
   ||:
Db 7 FFS 9

RESULT 39
Tl2312
cytochrome-c oxidase (EC 1.9.3.1) chain I - Sceloporus graciosus mitochondrion (fragment)
C:Species: mitochondrion Sceloporus graciosus
C>Date: 23-Jul-1999 #sequence_revision 23-Jul-1999 #text_change 22-Oct-1999
C:Accession: Tl2312
R:Schulte, J.A.; Macey, J.R.; Larson, A.; Papenfuss, T.J.
Mol. Phylogenet. Evol. 10, 367-376, 1998
A:Title: Molecular tests of phylogenetic taxonomies: A general procedure and example
A:Reference number: Z17488; MUID:99162288
A:Accession: Tl2312
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <SCH>
A:Cross-references: EMBL:AF049860; NID:g4105738; PID:g4105741; PIDN:AAD02523.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match      23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
   ||:
Db 7 FFS 9

RESULT 40
Tl2316
cytochrome-c oxidase (EC 1.9.3.1) chain I - Uma scoparia mitochondrion (fragment)
C:Species: mitochondrion Uma scoparia
C>Date: 23-Jul-1999 #sequence_revision 23-Jul-1999 #text_change 22-Oct-1999
C:Accession: Tl2316
R:Schulte, J.A.; Macey, J.R.; Larson, A.; Papenfuss, T.J.
Mol. Phylogenet. Evol. 10, 367-376, 1998
A:Title: Molecular tests of phylogenetic taxonomies: A general procedure and example
A:Reference number: Z17488; MUID:99162288
A:Accession: Tl2316
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <SCH>
A:Cross-references: EMBL:AF049861; NID:g4105742; PID:g4105745; PIDN:AAD02526.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match      23.6%; Score 13; DB 2; Length 10;

```

Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 41

T12321
Cytochrome-c oxidase (EC 1.9.3.1) chain I - Uta stansburiana mitochondrion (fragment)
C:Species: mitochondrion Uta stansburiana
C:Date: 23-Jul-1999 #sequence_revision 23-Jul-1999 #text_change 22-Oct-1999
C:Accession: T12321
R:Schulte, J.A.; Macey, J.R.; Larson, A.; Papenfuss, T.J.
Mol. Phylogenet. Evol. 10, 367-376, 1998
A:Title: Molecular tests of phylogenetic taxonomies: A general procedure and example used
A:Reference number: 217488; MUID:99162288
A:Accession: T12321
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <SCH>
A:Cross-references: EMBL:AF049863; NID:g4105750; PID:g4105753; PIDN:AAD02532.1
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 42

T14219
Cytochrome-c oxidase (EC 1.9.3.1) chain I - Xenosaurus grandis mitochondrion (fragment)
C:Species: mitochondrion Xenosaurus grandis
C:Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 21-Jul-2000
C:Accession: T14219
R:Macey, J.R.; Larson, A.; Ananjeva, N.B.; Fang, Z.; Papenfuss, T.J.
Mol. Biol. Evol. 14, 91-104, 1997
A:Title: Two novel gene orders and the role of light-strand replication in rearrangement
A:Reference number: 217789; MUID:97153826
A:Accession: T14219
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-10 <MAC>
A:Cross-references: EMBL:U71333; NID:g5739536; PIDN:AAC62821.1; PID:g1753275
C:Genetics:
A:Genome: mitochondrion
A:Note: COI
C:Keywords: mitochondrion; oxidoreductase

Query Match 23.6%; Score 13; DB 2; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.7e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 43

JQ1273
neuropeptide Antho-KAamide - sea anemone (Anthopleura elegantissima)
C:Species: Anthopleura elegantissima
C:Date: 31-Mar-1992 #sequence_revision 04-Dec-1992 #text_change 08-Dec-1995
C:Accession: JQ1273
R:Nothacker, H.P.; Rinehart, K.L.; Grimmelikhuijzen, C.J.P.

Biochem. Biophys. Res. Commun. 179, 1205-1211, 1991
A:Title: Isolation of L-3-phenyllactyl-Phe-Lys-Ala-NH2 (Antho-KAamide), a novel neuro
A:Reference number: JQ1273; MUID:92028852
A:Accession: JQ1273
A:Molecule type: protein
A:Residues: 1-4 <NOT>

C:Comment: The carboxyl-terminal amide probably arises from cleavage of a following g
C:Keywords: amidated carboxyl end; neuropeptide; phenyllactylation
F.1/Modified site: L-3-phenyllactic acid (Phe) #status experimental
F.4/Modified site: amidated carboxyl end (Ala) #status experimental

Query Match 21.8%; Score 12; DB 2; Length 4;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FF 8
||:
Db 1 FF 2

RESULT 44

A32480
achatin-I - giant African snail
N:Contains: achatin-II
C:Species: Achatina fulica (giant African snail)
C:Date: 12-Oct-1989 #sequence_revision 12-Oct-1989 #text_change 17-Mar-1999
C:Accession: A32480
R:Kamatani, Y.; Minakata, H.; Kenny, P.T.M.; Iwashita, T.; Watanabe, K.; Funase, K.;
Biochem. Biophys. Res. Commun. 160, 1015-1020, 1989
A:Title: Achatin-I, an endogenous neuroexcitatory tetrapeptide from Achatina fulica f
A:Reference number: A32480; MUID:89273551
A:Accession: A32480
A:Molecule type: protein
A:Residues: 1-4 <KAN>

A:Note: stereochemistry of the active form confirmed by chemical synthesis
R:Ishida, T.; In, Y.; Inoue, M.; Yasuda-Kamatani, Y.; Minakata, H.; Iwashita, T.; Nom
FEBS Lett. 307, 253-256, 1992
A:Title: Effect of the D-Phe(2) residue on molecular conformation of an endogenous ne
(H-Gly-Phe-Ala-Asp-OH).

A:Reference number: A44691; MUID:92354723
A:Contents: annotation; X-ray crystallography, 0.85 angstroms
A:Note: achatin-II has L-phenylalanine
C:Keywords: D-amino acid
F.2/Modified site: D-phenylalanine (Phe) #status experimental

Query Match 21.8%; Score 12; DB 2; Length 4;
Best Local Similarity 66.7%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
||:
Db 2 FAD 4

RESULT 45

A60986
N-formyl oligopeptide - Escherichia coli (fragment)
C:Species: Escherichia coli
C:Date: 31-Dec-1993 #sequence_revision 31-Dec-1993 #text_change 31-Dec-1993
C:Accession: A60986
R:Broom, M.F.; Mellor, D.M.; Chadwick, V.S.
Experientia 45, 1097-1099, 1989
A:Title: Purification and amino acid sequencing of naturally occurring N-formyl-methi
A:Reference number: A60986; MUID:90092408
A:Accession: A60986
A:Molecule type: protein
A:Residues: 1-6 <BRQ>

C:Comment: This hexapeptide was the longest of several N-formyl oligopeptides reporte
F.1/Modified site: N-formylmethionine #status experimental

Query Match 21.8%; Score 12; DB 2; Length 6;
Best Local Similarity 66.7%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 5 LVF 7
:||
Db 1 MVF 3

RESULT 46

159142
platelet-derived growth factor B chain - mouse (fragment)
C:Species: Mus musculus (house mouse)
C>Date: 02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change 05-Nov-1999
C:Accession: 159142
R:Pech, M.; Gazit, A.; Arnstein, P.; Aaronson, S.A.
Proc. Natl. Acad. Sci. U.S.A. 86, 2693-2697, 1989
A:Title: Generation of fibrosarcomas in vivo by a retrovirus that expressed the normal B cell myeloma protein gene
A:Reference number: 159142; MUID:89202393
A:Accession: 159142
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: mRNA
A:Residues: 1-6 <RES>
A:Cross-references: GB:M26180; NID:g516624; PIDN:AAA39905.1; PID:g516625

Query Match 21.8%; Score 12; DB 2; Length 6;
Best Local Similarity 66.7%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 5 LVF 7
:||
Db 1 MVF 3

RESULT 47

A43129
neuropeptide GNFRamide - tapeworm (Moniezia expansa)
C:Species: Moniezia expansa
C>Date: 10-Nov-1997 #sequence_revision 14-Nov-1997 #text_change 14-Nov-1997
C:Accession: A43129
R:Maule, A.; Shaw, C.; Halton, D.; Thim, L.
Biochem. Biophys. Res. Commun. 193, 1054-1060, 1993
A:Title: GNFRamide: A novel FMRFamide-immunoreactive peptide isolated from the sheep tapeworm Moniezia expansa
A:Reference number: A43129; MUID:93312289
A:Accession: A43129
A:Molecule type: protein
A:Residues: 1-6 <MAU>
C:Keywords: amidated carboxyl end; neuropeptide
F:6/Modified site: amidated carboxyl end (Phe) #status predicted

Query Match 21.8%; Score 12; DB 2; Length 6;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
||
Db 3 FF 4

RESULT 48

PT0246
Ig heavy chain CRD3 region (clone 2-103D) - human (fragment)
C:Species: Homo sapiens (man)
C>Date: 30-Sep-1993 #sequence_revision 30-Sep-1993 #text_change 16-Aug-1996
C:Accession: PT0246
R:Yamada, M.; Wasserman, R.; Reichard, B.A.; Shane, S.; Caton, A.J.; Rovera, G.
J. Exp. Med. 173, 395-407, 1991
A:Title: Preferential utilization of specific immunoglobulin heavy chain diversity and CDR3 hypervariability in the generation of the human antibody repertoire
A:Reference number: PT0246; MUID:91108337
A:Accession: PT0246
A:Molecule type: DNA
A:Residues: 1-7 <YAM>
A:Experimental source: B lymphocyte
C:Keywords: heterotrimer; immunoglobulin

Query Match 21.8%; Score 12; DB 2; Length 7;

Best Local Similarity 50.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HQKL 5
:|:
Db 1 HEVL 4

RESULT 49

I46868
alpha-myosin heavy chain - rabbit (fragment)
C:Species: Oryctolagus cuniculus (domestic rabbit)
C>Date: 14-Feb-1997 #sequence_revision 14-Feb-1997 #text_change 05-Nov-1999
C:Accession: I46868
R:Friedman, D.J.; Umeda, P.K.; Sinha, A.M.; Hsu, H.
Proc. Natl. Acad. Sci. U.S.A. 81, 3044-3048, 1984
A:Title: Characterization of genomic clones specifying rabbit alpha- and beta-ventricular myosin heavy chain genes
A:Reference number: I46868; MUID:84221901
A:Accession: I46868
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-7 <PRI>
A:Cross-references: GB:K01698; NID:g165538; PIDN:AAA31415.1; PID:g165539

Query Match 21.8%; Score 12; DB 2; Length 7;
Best Local Similarity 66.7%; Pred. No. 2.8e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QKL 5
:|:
Db 1 QKM 3

RESULT 50

T13818
cytochrome oxidase subunit I - Atlantic hagfish mitochondrion (fragment)
C:Species: mitochondrion Myxine glutinosa (Atlantic hagfish)
C>Date: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 21-Jul-2000
C:Accession: T13818
R:Delarabre, C.; Barriol, V.; Tillier, S.; Janvier, P.; Gachelin, G.
Mol. Biol. Evol. 14, 807-813, 1997
A:Title: The main features of the craniate mitochondrial DNA between the ND1 and the ND2 genes
A:Reference number: Z17775; MUID:97398704
A:Accession: T13818
A:Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-8

Query Match 21.8%; Score 12; DB 2; Length 8;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
||
Db 7 FF 8

Search completed: October 29, 2002, 09:38:49
Job time : 15 secs

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:31:37 ; Search time 11 Seconds
(without alignments)
35.200 Million cell updates/sec

Title: US-09-724-842a-27

Perfect score: 55

Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 105224 seqs, 38719550 residues

Total number of hits satisfying chosen parameters: 349

Minimum DB seq length: 0

Maximum DB seq length: 10

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 50 summaries

Database : SwlssProt_40:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Query Match %	Score	Length	DB ID	Description
1	18	32.7	10	1 COXM_RAT	P80431 rattus norv
2	16	29.1	5	1 RE11_LITRU	P82070 litoria rub
3	16	29.1	5	1 RE21_LITRU	P82071 litoria rub
4	16	29.1	10	1 PAP1_PARMA	P81863 pardachirus
5	15	27.3	5	1 UC22_MAIZE	P80628 zea mays (m
6	15	27.3	7	1 FAR2_ASCSU	P31890 ascaris suu
7	15	27.3	10	1 RCA_PINPS	P81084 pinus pinas
8	14	25.5	7	1 CCF1_ENTFA	P20104 enterococcu
9	14	25.5	8	1 CPD1_ENTFA	P13269 enterococcu
10	14	25.5	9	1 SAMP_MUSCA	P19095 mustelus ca
11	14	25.5	10	1 COXK_ONCMY	P80332 oncorhynch
12	13	23.6	8	1 UPAA_HUMAN	P30096 homo sapien
13	13	23.6	10	1 COXA_ONCMY	P80328 oncorhynch
14	13	23.6	10	1 COXO_RAT	P80432 rattus norv
15	12	21.8	4	1 ACH1_ACHFU	P35904 achatina fu
16	12	21.8	4	1 FFKA_ATEL	P58705 anthopleura
17	12	21.8	5	1 PAP2_PARMA	P81864 pardachirus
18	12	21.8	5	1 RE31_LITRU	P82072 litoria rub
19	12	21.8	5	1 RE32_LITRU	P82073 litoria rub
20	12	21.8	6	1 FARP_MONEX	P41966 moniezia ex
21	12	21.8	8	1 B44K_FORGI	P81886 porphyromon
22	12	21.8	9	1 FIBB_ERYPA	P19346 erythrocebu
23	12	21.8	10	1 COXO_THUOB	P80982 thunnus obe
24	12	21.8	10	1 FARP_MANSE	P18523 manduca sex
25	12	21.8	10	1 FARP_MYTED	P42560 mytilus edu
26	12	21.8	10	1 FIBB_CERSI	P14537 ceratotheri
27	12	21.8	10	1 MOSQ_CLYJA	P19962 clypeaster
28	12	21.8	10	1 TRNK_PIG	P01292 sus scrofa
29	12	21.8	10	1 TKU2_UREUN	P40752 urechis uni
30	12	21.8	10	1 TPIS_NICPL	P19118 nicotiana p
31	12	21.8	10	1 TRF6_LEUMA	P81738 leucophaea
32	12	21.8	10	1 TRP7_LEUMA	P81739 leucophaea
33	11	20.0	7	1 CHOX_ALCSP	P16101 alcaligenes

34	11	20.0	7	1 HV7_PIG	P01153 sus scrofa
35	11	20.0	7	1 UFO3_MOUSE	P38641 mus musculus
36	11	20.0	8	1 AKH_TABAT	P14595 tabanus atr
37	11	20.0	8	1 HTF2_PERAM	P04549 periplaneta
38	11	20.0	9	1 FAR5_PANRE	P82661 panagrellus
39	11	20.0	9	1 ULAK_MOUSE	P99031 mus musculus
40	11	20.0	10	1 BPP2_BOTIN	P30422 bothrops in
41	11	20.0	10	1 FAR6_PANRE	P82660 panagrellus
42	11	20.0	10	1 GON1_PETWA	P04378 petromyzon
43	11	20.0	10	1 HTF2_CARMO	P11385 carausius m
44	11	20.0	10	1 HTF_HELZE	P16353 heliothis z
45	11	20.0	10	1 HTF_TABAT	P14596 tabanus atr
46	11	20.0	10	1 Q2OB_COMTE	P80465 comamonas t
47	10	18.2	5	1 BPP7_BOTIN	P30425 bothrops in
48	10	18.2	8	1 ACL_THUAL	P18691 thunnus alb
49	10	18.2	8	1 ALL1_CYPDO	P82152 cydia pomon
50	10	18.2	8	1 LCR8_LEUMA	P19990 leucophaea

ALIGNMENTS

RESULT 1					
ID	COXM_RAT	STANDARD;	PRT;	10 AA.	
AC	P80431;				
DT	01-NOV-1995 (Rel. 32, Created)				
DT	01-NOV-1995 (Rel. 32, Last sequence update)				
DT	01-MAR-2002 (Rel. 41, Last annotation update)				
DE	Cytochrome c oxidase polypeptide VIIb, mitochondrial (EC 1.9.3.1) (Fragment).				
GN	COX7B.				
OS	Rattus norvegicus (Rat).				
OC	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;				
OC	Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.				
OX	NCBI_TaxID=10116;				
RN	[1]				
RP	SEQUENCE.				
RC	STRAIN=WISTAR; TISSUE=Liver;				
RX	MEDLINE=95324529; PubMed=7601105;				
RA	Schaeffer H., Noack H., Halangk W., Brandt U., von Jagow G.;				
RT	"cytochrome-c oxidase in developing rat heart. Enzymic properties and				
RT	amino-terminal sequences suggest identity of the fetal heart and the				
RL	adult liver isoform."				
RL	Eur. J. Biochem. 230:235-241(1995).				
CC	-!- FUNCTION: THIS PROTEIN IS ONE OF THE NUCLEAR-CODED POLYPEPTIDE				
CC	CHAINS OF CYTOCHROME C OXIDASE, THE TERMINAL OXIDASE IN				
CC	MITOCHONDRIAL ELECTRON TRANSPORT.				
CC	-!- CATALYTIC ACTIVITY: 4 ferrocytochrome c + O(2) = 4 ferricytochrome				
CC	c + 2 H(2)O.				
KW	Oxidoreductase; Mitochondrion.				
FT	NON_TER 10 10				
SQ	SEQUENCE 10 AA; 1210 MW; CFC70EB771A33326 CRC64;				
Query Match 32.7%; Score 18; DB 1; Length 10;					
Best Local Similarity 100.0%; Pred. No. 7.3e+02;					
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;					
Oy	2 HQK 4				
Db	2 HQK 4				
RESULT 2					
ID	RE11_LITRU	STANDARD;	PRT;	5 AA.	
AC	P82070;				
DT	01-MAR-2002 (Rel. 41, Created)				
DT	01-MAR-2002 (Rel. 41, Last sequence update)				
DT	01-MAR-2002 (Rel. 41, Last annotation update)				
DE	Rubellidin 1.1.				
OS	Litoria rubella (Desert tree frog).				
OC	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;				

DE Pardaxin I (EXI) (Fragment).
OS Pardachirus marmoratus (Red sea mooses sole).
OS Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC Acanthopterygia; Acanthopterygii; Percomorpha; Pleuronectiformes;
OC Soleioidi; Soleidae; Pardachirus.
OX NCBI_TaxID=31087;
RN [1]
RP SEQUENCE.
RP TIGSUE=Skin secretion;
RX MEDLINE=87057369; PubMed=3782138;
RA Lazarovici P., Primor N., Loew L.M.;
RT "Purification and pore-forming activity of two hydrophobic
RT polypeptides from the secretion of the Red sea mooses sole (Pardachirus
RT marmoratus).";
RL J. Biol. Chem. 261:16704-16713(1986).
CC -!- FUNCTION: EXHIBITS UNUSUAL SHARK REPELLENT AND SURFACTANT
CC PROPERTIES. FORMS VOLTAGE-DEPENDENT, ION-PERMEABLE CHANNELS IN
CC MEMBRANES. AT HIGH CONCENTRATION CAUSES CELL MEMBRANE LYSIS. SHOWN
CC TO BE 5-10 TIMES MORE TOXIC, CYTOLYTIC AND ACTIVE IN MEMBRANE PORE
CC FORMATION THAN PARADAXIN II.
CC -!- SUBUNIT: MONOMER. IN AQUEOUS SOLUTION EXISTS AS A TETRAMER.
CC -!- SUBCELLULAR LOCATION: Secreted.
CC -!- SIMILARITY: BELONGS TO THE PARADAXIN FAMILY.
FT Toxin.
KW NON_TER 10 10
KW SEQUENCE 10 AA; 1063 MW; D399C36760572DD9 CRC64;
SQ

Query Match 29.1%; Score 16; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.8e+03;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps

QY 7 FFA 9
DB |||
DB 2 FFA 4

RESULT 5
UC22_MAIZE
ID UC22_MAIZE STANDARD; PRT; 5 AA.
AC P80628;
DT 01-OCT-1996 (Rel. 34, Created)
DT 01-OCT-1996 (Rel. 34, Last sequence update)
DT 15-JUL-1999 (Rel. 38, Last annotation update)
DE Unknown protein from 2D-page of etioloated coleoptile (Spot 474)
DE (Fragment).
DE Zea mays (Maize).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; PACC clade;
OC Panicoideae; Andropogoneae; Zea.
OX NCBI_TaxID=4577;
RN [1]
RP SEQUENCE.
RP TISSUE=Coleoptile;
RA Touzet P., Riccardi F., Morin C., Damerval C., Huet J.-C.,
RA Perriollet J.-C., Zivy M., de Vienne D.;
RT "The maize two dimensional gel protein database: towards an integrated
RT genome analysis program.";
RL Theor. Appl. Genet. 93:997-1005(1996).
CC -!- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
CC PROTEIN IS: 6.1, ITS MW IS: 30.4 kDa.
CC Maize-2DPAGE: P80628; COLEOPTILE.
DR MaizeDB; 123954; -. 1
DR NON_TER 1 1
FT NON_TER 5 5
FT NON_TER 5 5
SQ SEQUENCE 5 AA; 654 MW; 72CB19C9C0300000 CRC64;

Query Match 27.3%; Score 15; DB 1; Length 5;
Best Local Similarity 66.7%; Pred. No. 1e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps

QY 6 VFF 8
DB :||

Db 1 IFF 3

RESULT 6

FAR2_ASCSU

ID FAR2_ASCSU STANDARD; PRT; 7 AA.
 AC P31890;
 DT 01-JUL-1993 (Rel. 26, Created)
 DT 01-JUL-1993 (Rel. 26, Last sequence update)
 DT 01-FEB-1996 (Rel. 33, Last annotation update)
 DE PWRamide-like neuropeptide AF2.
 OS Ascaris suum (big roundworm) (Ascaris lumbricoides), and
 OS Panagrellus redivivus.
 OC Eukaryota; Metazoa; Nematoda; Chromadorea; Ascaridida; Ascaridoidea;
 OC Ascarididae; Ascaris.
 OX NCBI_TaxID=6253, 6233;
 RN [1]
 RP SEQUENCE.
 RC SPECIES=A.suum;
 RX MEDLINE=93324431; PubMed=8332542;
 RA Cowden C., Stretton A.O.W.;
 RT "AF2, an Ascaris neuropeptide: isolation, sequence, and bioactivity.";
 RL Peptides 14:423-430(1993).
 RN [2]
 RP SEQUENCE.
 RC SPECIES=P.redivivus;
 RX MEDLINE=95060998; PubMed=7970891;
 RA Maule A.G., Shaw C., Bowman J.W.;
 RT "The FMRamide-like neuropeptide AF2 (Ascaris suum) is present in the
 RT free-living nematode, Panagrellus redivivus (Nematoda, Rhabditida).";
 RL Parasitology 109:351-356(1994).
 CC -|- FUNCTION: HAS EFFECTS ON MUSCLE TENSION.
 CC -|- TISSUE SPECIFICITY: FOUND IN THE NERVE CORDS AND A VARIETY OF
 CC GANGLIA PARTICULARLY IN THE ANTERIOR REGIONS.
 CC -|- SIMILARITY: BELONGS TO THE FARP (PWRAMIDE RELATED PEPTIDE)
 CC FAMILY.
 KW Neuropeptide; Amidation.
 FT MOD_RES 7;
 SQ SEQUENCE 7 AA; 992 MW; 69D4073B5B11E350 CRC64;

Query Match 27.3%; Score 15; DB 1; Length 7;
 Best Local Similarity 50.0%; Pred. No. 1e+05;
 Matches 3; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 2 HQLVF 7

I: I I

Db 2 HEVLR 7

RESULT 7

RCA_PINPS

ID RCA_PINPS STANDARD; PRT; 10 AA.
 AC P81084;
 DT 15-JUL-1998 (Rel. 36, Created)
 DT 15-JUL-1998 (Rel. 36, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Probable ribulose biphosphate carboxylase/oxygenase activase (RUBISCO
 DE activase) (RA) (Water stress responsive protein 4) (Fragment).
 OS Pinus pinaster (Maritime pine).
 OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
 OC Spermatophyta; Coniferopsida; Coniferales; Pinaceae; Pinus.
 OX NCBI_TaxID=71647;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Needle;
 RX MEDLINE=98418576; PubMed=9747804;
 RA Costa P., Bahrman N., Frigerio J.-M., Kremer A., Plomion C.;
 RT "Water-deficit-responsive proteins in maritime pine.";
 RL Plant Mol. Biol. 38:587-596(1998).
 RN [2]
 RP SEQUENCE.
 RC TISSUE=Needle;
 RX MEDLINE=99274088; PubMed=10344291;

RA Costa P., Plomion C., Bauw G., Dubos C., Bahrman N., Kremer A.,
 RA Frigerio J.-M., Plomion C.;
 RT "Separation and characterization of needle and xylem maritime pine
 RT proteins.";
 RL Electrophoresis 20:1098-1108(1999).
 CC -|- FUNCTION: ACTIVATION OF RUBISCO (RIBULOSE-1,5-BISPHOSPHATE
 CC CARBOXYLASE/OXYGENASE; EC 4.1.1.39) INVOLVES THE ATP-DEPENDENT
 CC CARBOXYLATION OF THE EPSILON-AMINO GROUP OF LYSINE LEADING TO A
 CC CARBAMATE STRUCTURE (BY SIMILARITY).
 CC -|- SUBCELLULAR LOCATION: Chloroplast stroma (By similarity).
 CC -|- INDUCTION: BY WATER STRESS.
 CC -|- SIMILARITY: BELONGS TO THE RUBISCO ACTIVASE FAMILY.
 KW Chloroplast; ATP-binding.
 FT NON_TER 1 1
 FT NON_TER 10 10
 SQ SEQUENCE 10 AA; 1171 MW; COA506D2C72B1EA6 CRC64;

Query Match 27.3%; Score 15; DB 1; Length 10;

Best Local Similarity 75.0%; Pred. No. 2.9e+03;

Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVF 7

: I I I

Db 5 ELVF 8

RESULT 8

CCFL_ENTFA

ID CCFL_ENTFA STANDARD; PRT; 7 AA.
 AC P20104;
 DT 01-FEB-1991 (Rel. 17, Created)
 DT 01-FEB-1991 (Rel. 17, Last sequence update)
 DT 01-FEB-1991 (Rel. 17, Last annotation update)
 DE Sex pheromone CCF10.
 OS Enterococcus faecalis (Streptococcus faecalis).
 OC Bacteria; Firmicutes; Bacillus/Clostridium group; Enterococcaceae;
 OC Enterococcus.
 OX NCBI_TaxID=1351;
 RN [1]
 RP SEQUENCE.
 RX MEDLINE=89008313; PubMed=3139658;
 RA Mori M., Sakagami Y., Ishii Y., Isogai A., Kitada C., Fujino M.,
 RA Adsit J.C., Dunn G.M., Suzuki A.;
 RT "Structure of ccf10, a peptide sex pheromone which induces
 RT conjugative transfer of the Streptococcus faecalis tetracycline
 RT resistance plasmid, pCF10.";
 RL J. Biol. Chem. 263:14574-14578(1988).
 CC -|- FUNCTION: CCF10 IS INVOLVED IN THE CONJUGATIVE TRANSFER OF THE
 CC HEMOLYSIN PLASMID PCF10.
 DR PIR; A30812; A30812.
 KW Pheromone.
 SQ SEQUENCE 7 AA; 790 MW; 72C9D2C731B2C740 CRC64;

Query Match 25.5%; Score 14; DB 1; Length 7;

Best Local Similarity 100.0%; Pred. No. 1e+05;

Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVF 7

I I I

Db 4 LVF 6

RESULT 9

CPDI_ENTFA

ID CPDI_ENTFA STANDARD; PRT; 8 AA.
 AC P13269;
 DT 01-JAN-1990 (Rel. 13, Created)
 DT 01-JAN-1990 (Rel. 13, Last sequence update)
 DT 01-FEB-1991 (Rel. 17, Last annotation update)
 DE Sex pheromone CPDI.
 OS Enterococcus faecalis (Streptococcus faecalis).
 OC Bacteria; Firmicutes; Bacillus/Clostridium group; Enterococcaceae;
 OC Enterococcus.

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OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE.
RX MEDLINE=85040388; PubMed=6436978;
RA Suzuki A., Mori M., Sagakami Y., Isogai A., Fujino M., Kitada C.,
RA Craig R.A., Clewell D.B.;
RT "Isolation and structure of bacterial sex pheromone, cpD1.";
RL Science 226:849-850(1984).
CC -!- FUNCTION: CPD1 IS INVOLVED IN THE CONJUGATIVE TRANSFER OF THE
CC BACTERIOCIN PLASMID PPd1.
KW Pheromone.
SQ SEQUENCE 8 AA; 913 MW; 8655B729C6820729 CRC64;

Query Match 25.5%; Score 14; DB 1; Length 8;
Best Local Similarity 75.0%; Pred. NO. 1e+05;
Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 5 LVFF 8
DB 2 LVNF 5

RESULT 10
SAMP_MUSCA STANDARD; PRT; 9 AA.
ID SAMP_MUSCA
AC P19095;
DT 01-NOV-1990 (Rel. 16, Created)
DT 01-NOV-1990 (Rel. 16, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Serum amyloid P-component (SAP) (Fragment).
OS Mustelus canis (Smooth dogfish).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Chondrichthyes;
OC Elasmobranchii; Galeomorphii; Galeoidea; Carcharhiniformes; Triakidae;
OC Mustelus.
OX NCBI_TaxID=7812;
RN [1]
RP SEQUENCE.
RX MEDLINE=83160932; PubMed=6403520;
RA Robey F.A., Tanaka T., Liu T.-Y.;
RT "Isolation and characterization of two major serum proteins from the
RT dogfish, Mustelus canis, C-reactive protein and amyloid P
RT component.";
RL J. Biol. Chem. 258:3889-3894(1983).
CC -!- SUBUNIT: HOMOPENTAMER. PENTAXIN (OR PENTRAXIN) HAVE A DISCOID
CC ARRANGEMENT OF 5 NONCOVALENTLY BOUND SUBUNITS.
CC -!- DISEASE: SAP IS A PRECURSOR OF AMYLOID COMPONENT P WHICH IS FOUND
CC IN BASEMENT MEMBRANE AND ASSOCIATED WITH AMYLOID DEPOSITS.
CC -!- SIMILARITY: BELONGS TO THE PENTAXIN FAMILY.
DR PIR; B20569; B20569.
DR InterPro; IPR001759; Pentaxin.
DR PROSITE; PS00289; PENTAXIN; PARTIAL.
KW Lectin; Amyloid; Glycoprotein; Plasma; Pentaxin.
FT DOMAIN 1 >9
FT NON_TER 9
FT SEQUENCE 9 AA; 965 MW; D05B5735B386769 CRC64;

Query Match 25.5%; Score 14; DB 1; Length 9;
Best Local Similarity 40.0%; Pred. NO. 1e+05;
Matches 2; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 3 OKLVF 7
DB 5 KSLIF 9

RESULT 11
COXK_ONCMY STANDARD; PRT; 10 AA.
ID COXK_ONCMY
AC P80332;
DT 01-OCT-1994 (Rel. 30, Created)
DT 01-OCT-1994 (Rel. 30, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Cytochrome c oxidase polypeptide VIIa-heart (EC 1.9.3.1) (Fragment).
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OS Oncorhynchus mykiss (Rainbow trout) (Salmo gairdneri).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei;
OC Protacanthopterygii; Salmoniformes; Salmonidae; Oncorhynchus.
OX NCBI_TaxID=8022;
RN [1]
RP SEQUENCE.
RX TISSUE=Heart;
RX MEDLINE=94237150; PubMed=8181469;
RA Freund R., Kadenbach B.;
RT "Identification of tissue-specific isoforms for subunits Vb and VIIa
RT of cytochrome c oxidase isolated from rainbow trout.";
RL Eur. J. Biochem. 221:1111-1116(1994).
CC -!- FUNCTION: THIS PROTEIN IS ONE OF THE NUCLEAR-CODED POLYPEPTIDE
CC CHAINS OF CYTOCHROME C OXIDASE, THE TERMINAL OXIDASE IN
CC MITOCHONDRIAL ELECTRON TRANSPORT.
CC -!- CATALYTIC ACTIVITY: 4 ferrocytochrome c + O(2) -> 4 ferricytochrome
CC c + 2 H(2)O.
CC -!- SUBCELLULAR LOCATION: Mitochondrial inner membrane.
CC -!- SIMILARITY: BELONGS TO THE CYTOCHROME C OXIDASE VIIA FAMILY.
DR PIR; S43631; S43631.
KW Oxidoreductase; Inner membrane; Mitochondrion.
FT NON_TER 10
FT SEQUENCE 10 AA; 1174 MW; 4C8D81CAFAF772C3 CRC64;

Query Match 25.5%; Score 14; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. NO. 4.5e+03;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 QKL 5
DB 8 QKL 10

RESULT 12
UPAA_HUMAN STANDARD; PRT; 8 AA.
ID UPAA_HUMAN
AC P30096;
DT 01-APR-1993 (Rel. 25, Created)
DT 01-APR-1993 (Rel. 25, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Unknown protein from 2D-page of plasma (Spot 36) (Fragment).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE.
RX TISSUE=Plasma;
RX MEDLINE=93092937; PubMed=1459097;
RA Hughes G.J., Frutiger S., Paquet N., Ravier F., Pasquali C.,
RA Sanchez J.-C., James R., Tissot J.-D., Bjellqvist B.,
RA Hochstrasser D.F.;
RT "Plasma protein map: an update by microsequencing.";
RL Electrophoresis 13:707-714(1992).
CC -!- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
CC PROTEIN IS: 7, ITS MW IS: 12 kDa.
DR SWISS-2DPAGE; P30096; HUMAN.
FT NON_TER 1
FT VARIANT 5 1 F -> P.
FT NON_TER 5 5 /FTid=VAR_000004.
FT SEQUENCE 8 AA; 909 MW; 86677B59D1A72042 CRC64;

Query Match 23.6%; Score 13; DB 1; Length 8;
Best Local Similarity 50.0%; Pred. NO. 1e+05;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 5 LVFF 8
DB 3 LTFY 6
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RESULT 13
COXA_ONCMY
ID COXA_ONCMY STANDARD; PRT; 10 AA.
AC P80328;
DT 01-OCT-1994 (Rel. 30, Created)
DT 01-OCT-1994 (Rel. 30, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Cytochrome c oxidase polypeptide VA (EC 1.9.3.1) (Fragment).
OS Oncorhynchus mykiss (Rainbow trout) (Salmo gairdneri).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei;
OC Protacanthopterygii; Salmoniformes; Salmonidae; Oncorhynchus.
OX NCBI_TaxID=8022;
RN [1]
RP SEQUENCE.
RC TISSUE=Liver;
RX MEDLINE=94237150; PubMed=8181469;
RA Freund R., Kadenbach B.;
RT "Identification of tissue-specific isoforms for subunits Vb and VIa
of cytochrome c oxidase isolated from rainbow trout.";
RL Eur. J. Biochem. 221:1111-1116(1994).
CC -!- FUNCTION: THIS IS THE HEME A-CONTAINING CHAIN OF CYTOCHROME C
OXIDASE, THE TERMINAL OXIDASE IN MITOCHONDRIAL ELECTRON TRANSPORT.
CC -!- CATALYTIC ACTIVITY: 4 ferrocytochrome c + O(2) = 4 ferricytochrome
c + 2 H(2)O.
CC -!- SUBCELLULAR LOCATION: Mitochondrial inner membrane.
CC -!- SIMILARITY: BELONGS TO THE CYTOCHROME C OXIDASE VA FAMILY.
DR PIR: S43625; S43625.
KW Oxidoreductase; Heme; Mitochondrion; Inner membrane.
FT NON_TER 10
FT SEQUENCE 10 AA; 1144 MW; C535C5B1AB02C33D CRC64;

Query Match 23.6%; Score 13; DB 1; Length 10;
Best Local Similarity 50.0%; Pred. No. 7e+03;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HOKL 5
| :
DB 2 HAKV 5

RESULT 14
COXO_RAT
ID COXO_RAT STANDARD; PRT; 10 AA.
AC P80432;
DT 01-NOV-1995 (Rel. 32, Created)
DT 01-NOV-1995 (Rel. 32, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Cytochrome c oxidase polypeptide VIIC, mitochondrial (EC 1.9.3.1)
DE (VIIIA) (Fragment).
GN COX7C OR COX7C1.
OS Rattus norvegicus (Rat).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX NCBI_TaxID=10116;
RN [1]
RP SEQUENCE.
RC STRAIN=WISTAR; TISSUE=Liver, and Heart;
RX MEDLINE=95324529; PubMed=7601105;
RA Schaeffer H., Noack H., Halangk W., Brandt U., von Jagow G.;
RT "Cytochrome-c oxidase in developing rat heart. Enzymic properties and
amino-terminal sequences suggest identity of the fetal heart and the
adult liver isoform".
RL Eur. J. Biochem. 230:235-241(1995).
CC -!- FUNCTION: THIS PROTEIN IS ONE OF THE NUCLEAR-CODED POLYPEPTIDE
CHAINS OF CYTOCHROME C OXIDASE, THE TERMINAL OXIDASE IN
MITOCHONDRIAL ELECTRON TRANSPORT.
CC -!- CATALYTIC ACTIVITY: 4 ferrocytochrome c + O(2) = 4 ferricytochrome
c + 2 H(2)O.
CC -!- SIMILARITY: BELONGS TO THE CYTOCHROME C OXIDASE VIIC FAMILY.
KW Oxidoreductase; Mitochondrion.
FT NON_TER 10
FT SEQUENCE 10 AA; 1117 MW; 126DE767687B1B0B CRC64;

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Query Match 23.6%; Score 13; DB 1; Length 10;
Best Local Similarity 25.0%; Pred. No. 7e+03;
Matches 1; Conservative 3; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQK 4
| : :
DB 2 HVEE 5

RESULT 15
ACH1_ACHFV
ID ACH1_ACHFV STANDARD; PRT; 4 AA.
AC P35904;
DT 01-JUN-1994 (Rel. 29, Created)
DT 01-JUN-1994 (Rel. 29, Last sequence update)
DT 15-JUL-1998 (Rel. 36, Last annotation update)
DE Achatin-I.
OS Achatina fulica (Giant African snail).
OC Eukaryota; Metazoa; Mollusca; Gastropoda; Pulmonata; Stylommatophora;
OC Achatinacea; Achatinidae; Achatina.
OX NCBI_TaxID=6530;
RN [1]
RP SEQUENCE, CHARACTERIZATION, AND SYNTHESIS.
RC STRAIN=FERUSSAC; TISSUE=Ganglion;
RX MEDLINE=89273551; PubMed=2597281;
RA Kanatani Y., Minakata H., Kenny P.T.M., Iwashita T., Watanabe K.,
Furuse K., Sun X.P., Yongsiri A., Kim K.H., Novales-Li P.,
RA Novales E.T., Kanai C.G., Takeuchi H., Nomoto K.;
RT "Achatin-I, an endogenous neuroexcitatory tetrapeptide from Achatina
fulica Ferussac containing a D-amino acid residue.";
RL Biochem. Biophys. Res. Commun. 160:1015-1020(1989).
RN [2]
RP CHARACTERIZATION.
RC STRAIN=FERUSSAC; TISSUE=Heart atrium;
RX MEDLINE=91264856; PubMed=1675568;
RA Fujimoto K., Kubota I., Yasuda-Kamatani Y., Minakata H., Nomoto K.,
RA Yoshida M., Harada A., Muneoka Y., Kobayashi M.;
RT "Purification of achatin-I from the atria of the African giant snail,
Achatina fulica, and its possible function.";
RL Biochem. Biophys. Res. Commun. 177:847-853(1991).
RN [3]
RP X-RAY CRYSTALLOGRAPHY.
RX MEDLINE=93014529; PubMed=1399265;
RA Ishida T., In Y., Doi M., Inoue M., Yasuda-Kamatani Y., Minakata H.,
RA Iwashita T., Nomoto K.;
RT "Crystal structure and molecular conformation of achatin-I
(H-Gly-D-Phe-Ala-Asp-OH), an endogenous neuropeptide containing a
D-amino acid residue.";
RL Int. J. Pept. Protein Res. 39:258-264(1992).
CC -!- FUNCTION: NEUROEXCITATORY PEPTIDE; INCREASES THE IMPULSE FREQUENCY
AND PRODUCES A SPIKE BROADENING OF THE IDENTIFIED HEART EXCITATORY
NEURON (PON); ALSO ENHANCES THE AMPLITUDE AND FREQUENCY OF THE
HEART BEAT. HAS ALSO AN EFFECT ON SEVERAL OTHER MUSCLES.
DR PIR: A32480; A32480.
KW Hormone; D-amino acid.
FT MOD_RES 2
FT SEQUENCE 4 AA; 408 MW; 6AADD9C810000000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 4;
Best Local Similarity 66.7%; Pred. No. 1e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
| :
DB 2 FAD 4

RESULT 16
FFKA_ATEL
ID FFKA_ATEL STANDARD; PRT; 4 AA.
AC P58705;
DT 01-MAR-2002 (Rel. 41, Created)

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DT 01-MAR-2002 (Rel. 41, Last sequence update)
DE 01-MAR-2002 (Rel. 41, Last annotation update)
DE Antho-KAamide.
OS Anthopleura elegantissima (Sea anemone).
OC Eukaryota; Metazoa; Cnidaria; Anthozoa; Zoantharia; Actiniaria;
OC Nynanthaeae; Actiniidae; Anthopleura.
OX NCBI_TaxID=6110;
RN [1]
RN SEQUENCE.
RX PubMed=1681803;
RA Notherack H.-P., Rinehart K.L. Jr., Grimmelikhuijzen C.J.P.;
RT "Isolation of L-3-phenyllactyl-Phe-Lys-Ala-NH2 (Antho-KAamide), a
RT novel neuropeptide from sea anemones.";
RL Biochem. Biophys. Res. Commun. 179:1205-1211(1991).
RN [2]
RN FUNCTION.
RX PubMed=8397415;
RA McFarlane I.D., Hudman D., Notherack H.-P., Grimmelikhuijzen C.J.P.;
RT "The expansion behaviour of sea anemones may be coordinated by two
RT inhibitory neuropeptides, Antho-KAamide and Antho-Riamide.";
RL Proc. R. Soc. Lond., B, Biol. Sci. 253:183-188(1993).
CC -1- FUNCTION: Inhibits spontaneous contractions in several muscle
CC groups. May be involved in the expansion phase of feeding
CC behaviour in sea anemones.
CC -1- SUBCELLULAR LOCATION: Secreted.
CC -1- TISSUE SPECIFICITY: Neuron-specific.
KW Neuropeptide; Amidation.
FT MOD_RES 1 1 L-3-PHENYLLACTYL.
FT MOD_RES 4 4 AMIDATION.
SQ SEQUENCE 4 AA; 512 MW; 6DD339C9A0000000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 4;
Best Local Similarity 100.0%; Pred. No. le+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FF 8
||
Db 1 FF 2

RESULT 17
PAP2_PARMA
ID PAP2_PARMA STANDARD; PRT; 5 AA.
AC P81864;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DE 16-OCT-2001 (Rel. 40, Last annotation update)
DE Pardaxin II (PXII) (Fragment).
OS Pardachirus marmoratus (Red sea mores sole).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC Acanthomorpha; Acanthopterygii; Percormorpha; Pleuronectiformes;
OC Soleoidel; Soleidae; Pardachirus.
OX NCBI_TaxID=31087;
RN [1]
RN SEQUENCE.
RX MEDLINE=87057369; PubMed=3782138;
RA Lazarovici P., Primor N., Loew L.M.;
RT "Purification and pore-forming activity of two hydrophobic
RT polypeptides from the secretion of the Red sea mores sole (Pardachirus
RT marmoratus).";
RL J. Biol. Chem. 261:16704-16713(1986).
CC -1- FUNCTION: EXHIBITS UNUSUAL SHARK REPELLENT AND SURFACTANT
CC PROPERTIES. FORMS VOLTAGE-DEPENDENT, ION-PERMEABLE CHANNELS
CC IN MEMBRANES. AT HIGH CONCENTRATION CAUSES CELL MEMBRANE LYSIS.
CC -1- SUBUNIT: MONOMER. IN AQUEOUS SOLUTION EXISTS AS A TETRAMER.
CC -1- SUBCELLULAR LOCATION: Secreted.
CC -1- SIMILARITY: BELONGS TO THE PARDAXIN FAMILY.
KW Toxin.
FT NON_TER 5 5
SQ SEQUENCE 5 AA; 614 MW; 7769C9C8100000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 5;
Best Local Similarity 100.0%; Pred. No. le+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FF 8
||
Db 2 FF 3

RESULT 18
RE31_LITRU
ID RE31_LITRU STANDARD; PRT; 5 AA.
AC P82072;
DT 01-MAR-2002 (Rel. 41, Created)
DT 01-MAR-2002 (Rel. 41, Last sequence update)
DE 01-MAR-2002 (Rel. 41, Last annotation update)
DE Rubellidin 3.1.
OS Litoria rubella (Desert tree frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Neobatrachia; Bufonoidea; Hylidae;
OC Litoria.
OX NCBI_TaxID=104895;
RN [1]
RN SEQUENCE, AND MASS SPECTROMETRY.
RP TISSUE-Skin secretion;
RA Steinborner S.T., Wabnitz P.A., Waugh R.J., Bowie J.H., Gao C.,
RA Tyler M.J., Wallace J.C.;
RT "The structure of new peptides from the Australian red tree frog
RT 'Litoria rubella', the skin peptide profile as a probe for the study
RT of evolutionary trends of amphibians.";
RL Aust. J. Chem. 49:955-963(1996).
CC -1- FUNCTION: SHOW NEITHER NEUROPEPTIDE ACTIVITY NOR ANTIBIOTIC
CC ACTIVITY.
CC -1- TISSUE SPECIFICITY: SECRETED BY THE SKIN DORSAL GLANDS.
KW Amphibian skin; Amidation.
FT MOD_RES 5 5 AMIDATION.
SQ SEQUENCE 5 AA; 656 MW; 71A9C9CB10300000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 5;
Best Local Similarity 100.0%; Pred. No. le+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FF 8
||
Db 3 FF 4

RESULT 19
RE32_LITRU
ID RE32_LITRU STANDARD; PRT; 5 AA.
AC P82073;
DT 01-MAR-2002 (Rel. 41, Created)
DT 01-MAR-2002 (Rel. 41, Last sequence update)
DE 01-MAR-2002 (Rel. 41, Last annotation update)
DE Rubellidin 3.2.
OS Litoria rubella (Desert tree frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Neobatrachia; Bufonoidea; Hylidae;
OC Litoria.
OX NCBI_TaxID=104895;
RN [1]
RN SEQUENCE.
RP TISSUE-Skin secretion;
RA Wabnitz P.A., Bowie J.H., Tyler M.J., Wallace J.C.;
RT "Peptides from the skin glands of the Australian buzzing tree frog
RT Litoria electrica. Comparison with the skin peptides from Litoria
RT rubella.";
RL Aust. J. Chem. 52:0-0(1999).
CC -1- FUNCTION: SHOW NEITHER NEUROPEPTIDE ACTIVITY NOR ANTIBIOTIC
CC ACTIVITY.
CC -1- TISSUE SPECIFICITY: SECRETED BY THE SKIN DORSAL GLANDS.
KW Amphibian skin.

SO SEQUENCE 5 AA; 570 MW; 71A9C9C862A00000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 5;
Best Local Similarity 100.0%; Pred. No. 1e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 3 FF 4

RESULT 20

FARP_MONEX

ID FARP_MONEX STANDARD; PRT; 6 AA.

AC P41966;

DT 01-NOV-1995 (Rel. 32, Created)

DT 01-NOV-1995 (Rel. 32, Last sequence update)

DT 01-NOV-1997 (Rel. 35, Last annotation update)

DE FMRamide-like neuropeptide GNFRF-amide.

OS Monieza expansa (Sheep tapeworm).

OC Eukaryota; Metazoa; Platyhelminthes; Turbellarian Platyhelminths;

OC Rhabditophora; Eulecithophora; Revertospermata; Mediofusata;

OC Neodermata; Cestoda; Eucestoda; Cyclophyllidae; Anoplocephalidae;

OC Monieza.

OX NCBI_TaxID=28841;

RN [1]

RP SEQUENCE.

RA MEDLINE=93312289; PubMed=8323531;

RT Maule A.G., Shaw C., Halton D.W., Thim L.;

RT "GNFRamide: a novel FMRamide-immunoreactive peptide isolated from

the sheep tapeworm, Monieza expansa.";

RL Biochem. Biophys. Res. Commun. 193:1054-1060(1993).

CC -1- SIMILARITY: BELONGS TO THE FARP (FMRamide RELATED PEPTIDE)

CC FAMILY.

KW Neuropeptide; Amidation.

FT MOD_RES 6 6

SO SEQUENCE 6 AA; 787 MW; 69D409C9C4481000 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 6;
Best Local Similarity 100.0%; Pred. No. 1e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 3 FF 4

RESULT 21

B44K_PORGI

ID B44K_PORGI STANDARD; PRT; 8 AA.

AC P81886;

DT 30-MAY-2000 (Rel. 39, Created)

DT 30-MAY-2000 (Rel. 39, Last sequence update)

DT 30-MAY-2000 (Rel. 39, Last annotation update)

DE 44 kDa immunogenic protein (Fragment).

OS Porphyromonas gingivalis (Bacteroides gingivalis).

OC Bacteria; CFB group; Bacteroidetes; Bacteroidales; Porphyromonadaceae;

OC Porphyromonas.

OX NCBI_TaxID=837;

RN [1]

RP SEQUENCE.

RC STRAIN=VPB 3492;

RX MEDLINE=20198497; PubMed=10731616;

RA Norris J.M., Love D.N.;

RT "Serum antibody responses of cats to soluble whole cell antigens of

feline Porphyromonas gingivalis.";

RL Vet. Microbiol. 73:37-49(2000).

CC -1- SIMILARITY: TO P.GINGIVALIS HEMAGGLUTININ A.

KW Antigen.

FT NON_TER 8 8

SO SEQUENCE 8 AA; 989 MW; 9554540326CB476D CRC64;

Query Match

21.8%; Score 12; DB 1; Length 8;

Best Local Similarity 66.7%; Pred. No. 1e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HQK 4
Db 3 YOK 5

RESULT 22

FIBB_ERYPA

ID FIBB_ERYPA STANDARD; PRT; 9 AA.

AC P19346;

DT 01-NOV-1990 (Rel. 16, Created)

DT 01-NOV-1990 (Rel. 16, Last sequence update)

DT 01-MAR-2002 (Rel. 41, Last annotation update)

DE Fibrinogen beta chain (Contains: Fibrinopeptide B) (Fragment).

GN FGB.

OS Erythrocybus patas (Red guenon) (Hussar).

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

OC Mammalia; Eutheria; Primates; Catarrhini; Cercopitheciidae;

OC Cercopitheciinae; Erythrocybus.

OX NCBI_TaxID=9538;

RN [1]

RP SEQUENCE.

RX MEDLINE=85289140; PubMed=3928610;

RA Nakamura S., Takenaka O., Takahashi K.;

RT "Fibrinopeptides A and B of Japanese monkey (Macaca fuscata) and

patas monkey (Erythrocybus patas): their amino acid sequences,

restricted mutations, and a molecular phylogeny for macaques,

guenons, and baboons.";

RL J. Biochem. 97:1487-1492(1985).

CC -1- FUNCTION: FIBRINOGEN HAS A DOUBLE FUNCTION: YIELDING MONOMERS THAT

POLYMERIZE INTO FIBRIN AND ACTING AS A COFACTOR IN PLATELET

AGGREGATION.

CC -1- SUBUNIT: HEXAMER CONTAINING 2 SETS OF 3 NONIDENTICAL CHAINS

(ALPHA, BETA AND GAMMA), LINKED TO EACH OTHER BY DISULFIDE BONDS.

CC -1- MISCELLANEOUS: CONVERSION OF FIBRINOGEN TO FIBRIN IS TRIGGERED BY

THROMBIN, WHICH CLEAVES FIBRINOPEPTIDES A AND B FROM ALPHA & BETA

CHAINS, AND THUS EXPOSES THE N-TERMINAL POLYMERIZATION SITES

RESPONSIBLE FOR THE FORMATION OF THE SOFT CLOT.

DR PIR: D24180; D24180.

DR InterPro: IPR002181; Fibrinogen C.

DR PROSITE: PS00514; FIBRIN_AG_C_DOMAIN; PARTIAL.

KW Blood coagulation; Plasma.

FT PEPTIDE 1 9 FIBRINOPEPTIDE B.

FT NON_TER 9 9

SO SEQUENCE 9 AA; 1020 MW; 69FE7879C732CB1B CRC64;

Query Match

21.8%; Score 12; DB 1; Length 9;

Best Local Similarity 16.7%; Pred. No. 1e+05;
Matches 1; Conservative 5; Mismatches 0; Indels 0; Gaps 0;

Qy 2 HOKLVF 7
Db 1 NEEVLF 6

RESULT 23

COXO_THUOB

ID COXO_THUOB STANDARD; PRT; 10 AA.

AC P80982;

DT 01-NOV-1997 (Rel. 35, Created)

DT 01-NOV-1997 (Rel. 35, Last sequence update)

DT 16-OCT-2001 (Rel. 40, Last annotation update)

DE Cytochrome c oxidase polypeptide VIIC (EC 1.9.3.1) (Fragment).

OS Thunnus obesus (Bigeye tuna).

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;

OC Acanthomorpha; Acanthopterygii; Percomorpha; Perciformes; Scombroidei;

OC Scombridae; Thunnus.

OX NCBI_TaxID=8241;

RN [1]

RP SEQUENCE.

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RC TISSUE=Heart, and Liver;
RX MEDLINE=97454291; PubMed=9310366;
RA Arnold S., Lee I., Kim M., Song E., Linder D., Lottspeich F.,
RT Kadenbach B.;
RT "The subunit structure of cytochrome-c oxidase from tuna heart and
RL liver.";
RL Eur. J. Biochem. 248:99-103(1997).
CC -!- FUNCTION: THIS PROTEIN IS ONE OF THE NUCLEAR-CODED POLYPEPTIDE
CC CHAINS OF CYTOCHROME C OXIDASE, THE TERMINAL OXIDASE IN
CC MITOCHONDRIAL ELECTRON TRANSPORT.
CC -!- CATALYTIC ACTIVITY: 4 ferrocycochrome c + O(2) = 4 ferricytochrome
CC c + 2 H(2)O.
CC -!- SUBCELLULAR LOCATION: Mitochondrial inner membrane.
CC -!- SIMILARITY: BELONGS TO THE CYTOCHROME C OXIDASE VIIC FAMILY.
KW Oxidoreductase; Inner membrane; Mitochondrion.
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1059 MW; 126DE767687B1DCB CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 66.7%; Pred. No. 1.1e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
   :||
DB 3 YAE 5

RESULT 24
FARP_MANSE STANDARD; PRT; 10 AA.
AC P18523;
DT 01-NOV-1990 (Rel. 16, Created)
DT 01-FEB-1994 (Rel. 28, Last sequence update)
DT 01-NOV-1995 (Rel. 32, Last annotation update)
DE FMRFamide-like neuropeptide.
OS Manduca sexta (Tobacco hawkmoth) (Tobacco hornworm).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia;
OC Sphingioidea; Sphingidae; Sphinginae; Manduca.
OX NCBI_TaxID=7130;
RN [1]
RP SEQUENCE.
RX MEDLINE=91045350; PubMed=2235684;
RA Kingan T.G., Replow D.B., Phillips J.M., Riehm J.P., Rao K.R.,
RA Hildebrand J.G., Homberg U., Kammer A.E., Jardine I., Griffin P.R.,
RA Hunt D.F.;
RT "A new peptide in the FMRFamide family isolated from the CNS of the
RT hawkmoth, Manduca sexta.";
RL Peptides 11:849-856(1990).
CC -!- FUNCTION: INCREASES THE FORCE OF NEURALLY EVOKED CONTRACTIONS IN
CC THE MAJOR POWER-PRODUCING FLIGHT MUSCLES, THE DORSAL LONGITUDINAL
CC MUSCLES AND SO IS LIKELY TO PLAY A ROLE IN SUSTAINING OR PROMOTING
CC FLIGHT BEHAVIOR PATTERNS.
CC -!- SIMILARITY: BELONGS TO THE FARP (FMRFAMIDE RELATED PEPTIDE)
CC FAMILY.
DR PIR; A43977; A43977.
KW Amidation; Neuropeptide.
FT MOD_RES 1 1 PYRROLIDONE CARBOXYLIC ACID.
FT MOD_RES 10 10 AMIDATION.
SQ SEQUENCE 10 AA; 1247 MW; D3C4529D5B1F2D2 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 50.0%; Pred. No. 1.1e+04;
Matches 3; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 2 HOKLVF 7
   :||
DB 5 HSFLRF 10

RESULT 25
FARP_MYTED STANDARD; PRT; 10 AA.
ID FARP_MYTED

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AC P42560;
DT 01-NOV-1995 (Rel. 32, Created)
DT 01-NOV-1995 (Rel. 32, Last sequence update)
DE FMRFamide-like neuropeptide ALAGDHFRRF-amide.
OS Mytilus edulis (Blue mussel).
OC Eukaryota; Metazoa; Mollusca; Bivalvia; Pteriomorpha; Mytiloidea;
OC Mytiloidea; Mytilidae; Mytilus.
OX NCBI_TaxID=6550;
RN [1]
RP SEQUENCE.
RX MEDLINE=93047883; PubMed=1358534;
RA Walker R.J.;
RT "Neuroactive peptides with an RFamide or Famide carboxyl terminal.";
RL Comp. Biochem. Physiol. 102C:213-222(1992).
CC -!- SIMILARITY: BELONGS TO THE FARP (FMRFAMIDE RELATED PEPTIDE)
CC FAMILY.
KW Neuropeptide; Amidation.
FT MOD_RES 10 10 AMIDATION.
SQ SEQUENCE 10 AA; 1180 MW; C2F80CC9C1EAA87D CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FF 8
   :||
DB 7 FF 8

RESULT 26
FIBB_CERSI STANDARD; PRT; 10 AA.
ID FIBB_CERSI
AC P14537;
DT 01-JAN-1990 (Rel. 13, Created)
DT 01-JAN-1990 (Rel. 13, Last sequence update)
DT 01-MAR-2002 (Rel. 41, Last annotation update)
DE Fibrinogen beta chain [Contains: Fibrinopeptide B] (Fragment).
GN FGB.
OS Ceratotherium simum (White rhinoceros) (Square-lipped rhinoceros).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Perissodactyla; Rhinocerotidae; Ceratotherium.
OX NCBI_TaxID=9807;
RN [1]
RP SEQUENCE.
RA O'Neill P.B., Doolittle R.F.;
RT "Mammalian phylogeny based on fibrinopeptide amino acid sequences.";
RL Syst. Zool. 22:590-595(1973).
CC -!- FUNCTION: FIBRINOGEN HAS A DOUBLE FUNCTION: YIELDING MONOMERS THAT
CC POLYMERIZE INTO FIBRIN AND ACTING AS A COFACTOR IN PLATELET
CC AGGREGATION.
CC -!- SUBUNIT: HEXAMER CONTAINING 2 SETS OF 3 NONIDENTICAL CHAINS
CC (ALPHA, BETA AND GAMMA), LINKED TO EACH OTHER BY DISULFIDE BONDS.
CC -!- MISCELLANEOUS: CONVERSION OF FIBRINOGEN TO FIBRIN IS TRIGGERED BY
CC THROMBIN, WHICH CLEAVES FIBRINOPEPTIDES A AND B FROM ALPHA & BETA
CC CHAINS, AND THUS EXPOSES THE N-TERMINAL POLYMERIZATION SITES
CC RESPONSIBLE FOR THE FORMATION OF THE SOFT CLOT.
DR InterPro; IPR002181; Fibrinogen_C.
DR PROSITE; PS00514; FIBRIN_AG_C_DOMAIN; PARTIAL.
KW Blood coagulation; Plasma.
FT PEPTIDE 1 10 FIBRINOPEPTIDE B.
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1097 MW; 9402B2B2CDDDD33A CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 50.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQK 4
   :||
DB 1 HDDK 4

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RESULT 27

MOSQ_CLYJA
ID MOSQ_CLYJA STANDARD; PRT; 10 AA.
AC P19962;
DT 01-FEB-1991 (Rel. 17, Created)
DT 01-FEB-1991 (Rel. 17, Last sequence update)
DT 01-FEB-1991 (Rel. 17, Last annotation update)
DE [Gln-6]-mosact
OS Clypeaster japonicus (Sand dollar).
OC Eukaryota; Metazoa; Echinodermata; Eleutherozoa; Echinozoa;
OC Echinoidea; Euechinoidea; Gnathostomata; Clypeasteroidea;
OC Clypeasteridae; Clypeaster.
OX NCBI_TaxID=7644;
RN [1]
RP SEQUENCE.
RC TISSUE=Egg jelly;
RA Suzuki N., Kurita M., Yoshino K., Kajlura H., Nomura K., Yamaguchi M.;
RT "Purification and structure of mosact and its derivatives from the
RT egg jelly of the sea urchin Clypeaster japonicus.";
RL 2001. Sci. 4: 649-656(1987).
CC -1- FUNCTION: STIMULATES SPERM RESPIRATION AND MOTILITY.
DR PIR; JN0025; JN0025.
SQ SEQUENCE 10 AA; 1019 MW; 9AFB032456DDC5BA CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 50.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 3 QKLV 6
| |
Db 6 QNLI 9

RESULT 28

TNKK_PIG
ID TNKK_PIG STANDARD; PRT; 10 AA.
AC P01292;
DT 21-JUL-1986 (Rel. 01, Created)
DT 21-JUL-1986 (Rel. 01, Last sequence update)
DT 16-OCT-2001 (Rel. 40, Last annotation update)
DE Neurokinin B (NKB) (Neuromedin K).
GN TAC3 OR NKNB.
OS Sus scrofa (Pig), and
OS Rana ridibunda (Laughing frog) (Marsh frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
OX NCBI_TaxID=9823, 8406;
RN [1]
RP SEQUENCE.
RC SPECIES=Pig; TISSUE=Spinal cord;
RX MEDLINE=83282812; PubMed=6576785;
RA Kangawa K., Minamino N., Fukuda A., Matsuo H.;
RT "Neuromedin K: a novel mammalian tachykinin identified in porcine
RT spinal cord.";
RL Biochem. Biophys. Res. Commun. 114:533-540(1983).
RN [2]

SEQUENCE.
RC SPECIES=R. ridibunda; TISSUE=Brain;
RX MEDLINE=92044543; PubMed=1658233;
RA O'Harte F., Burcher E., Lovas S., Smith D.D., Vaudry H., Conlon J.M.;
RT "Ranakinin: a novel NK1 tachykinin receptor agonist isolated with
RT neurokinin B from the brain of the frog Rana ridibunda.";
RL J. Neurochem. 57:2086-2091(1991).
CC -1- FUNCTION: TACHYKININS ARE ACTIVE PEPTIDES WHICH EXCITE NEURONS,
CC EVOKE BEHAVIORAL RESPONSES, ARE POTENT VASODILATORS AND
CC SECRETAGOGUES, AND CONTRACT (DIRECTLY OR INDIRECTLY) MANY SMOOTH
CC MUSCLES.
CC -1- SUBCELLULAR LOCATION: Secreted.
CC PIR; A01560; SPGNK.
DR InterPro; IPR002040; Tachykinin.
DR PROSITE; PS00267; TACHYKININ; 1.
KW Tachykinin; Neuropeptide; Amidation.

FT MOD_RES 10 10 AMIDATION
SQ SEQUENCE 10 AA; 1211 MW; E1FA7C62C9C9CAAL CRC64;
Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 7 FF 8
| |
Db 5 FF 6
RESULT 29
TKU2_UREUN
ID TKU2_UREUN STANDARD; PRT; 10 AA.
AC P40752;
DT 01-FEB-1995 (Rel. 31, Created)
DT 01-FEB-1995 (Rel. 31, Last sequence update)
DT 01-NOV-1995 (Rel. 32, Last annotation update)
DE Urechis tachykinin II.
OS Urechis unicinctus.
OC Eukaryota; Metazoa; Echiura; Xenopneusta; Urechidae; Urechis.
OX NCBI_TaxID=6432;
RN [1]
RP SEQUENCE, AND SYNTHESIS.
RC TISSUE=Ventral nerve cord;
RX MEDLINE=93236558; PubMed=8476410;
RA Ikeda T., Minakata H., Nomoto K., Kubota I., Muneoka Y.;
RT "Two novel tachykinin-related neuropeptides in the echiuroid worm,
RT Urechis unicinctus.";
RL Biochem. Biophys. Res. Commun. 192:1-6(1993).
CC -1- FUNCTION: CONTRACTILE ACTION ON THE INNER CIRCULAR BODY-WALL
CC MUSCLE OF THE ANIMAL.
CC -1- SIMILARITY: SOME SIMILARITY TO TACHYKININS.
KW Tachykinin; Neuropeptide; Amidation.
FT MOD_RES 10 10 AMIDATION.
SQ SEQUENCE 10 AA; 984 MW; 3F58DD79C87698 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
| |
Db 6 FF 7

RESULT 30

TPIS_NICPL
ID TPIS_NICPL STANDARD; PRT; 10 AA.
AC P19118;
DT 01-NOV-1990 (Rel. 16, Created)
DT 01-NOV-1990 (Rel. 16, Last sequence update)
DT 01-FEB-1996 (Rel. 33, Last annotation update)
DE Triosephosphate isomerase, cytosolic (EC 5.3.1.1) (TIM) (Fragment).
OS Nicotiana glauca (Leadwort-leaved tobacco).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC Asteridae; euasterids I; Solanales; Solanaceae; Nicotiana.
OX NCBI_TaxID=4092;
RN [1]
RP SEQUENCE.
RA Bauw G., de Loose M., Inze D., van Montagu M., Vandekerckhove J.;
RT "Alterations in the phenotype of plant cells studied by NH2-terminal
RT amino acid-sequence analysis of proteins electrophoretically from two-
RT dimensional gel-separated total extracts.";
RL Proc. Natl. Acad. Sci. U.S.A. 84:4806-4810(1987).
CC -1- CATALYTIC ACTIVITY: D-glyceraldehyde 3-phosphate -> glyceralone
CC phosphate.
CC -1- PATHWAY: PLAYS AN IMPORTANT ROLE IN SEVERAL METABOLIC PATHWAYS.
CC -1- SUBUNIT: HOMODIMER.
CC -1- SUBCELLULAR LOCATION: Cytoplasmic (probable).
CC -1- MISCELLANEOUS: IN PLANTS, THERE ARE TWO TYPES OF TPIS, CYTOSOLIC

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CC -1- SIMILARITY: BELONGS TO THE TRIOSEPHOSPHATE ISOMERASE FAMILY.
DR PIR: A27617; A27617.
DR InterPro: IPR000652; Trioseph_isomerase.
DR Pfam: PF00121; TIM; 1.
DR PROSITE: PS00171; TIM; PARTIAL.
KW Isomerase; Glycolysis; Gluconeogenesis; Fatty acid biosynthesis;
KW Pentose shunt.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1140 MW; 80B9D37862C9C9D1 CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 4 FF 5

RESULT 31
TRP6_LEUMA
ID TRP6_LEUMA STANDARD; PRT; 10 AA.
AC P81738;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Tachykinin-related peptide 6 (LemTRP 6).
OS Leucophaea maderae (Madeira cockroach).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Orthopteroidea; Dictyoptera; Blattaria;
OC Blaberoidea; Blaberidae; Leucophaea.
RN NCBI_TaxID=6988;
[1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=Brain;
RX MEDLINE=97262666; PubMed=9114447;
RA Muren J.E., Naessel D.R.;
RT "Seven tachykinin-related peptides isolated from the brain of the
RT Madeira cockroach; evidence for tissue-specific expression of
RT isoforms."
RL Peptides 18:7-15(1997).
CC -1- FUNCTION: MYOACTIVE PEPTIDE. INCREASES THE AMPLITUDE AND FREQUENCY
CC OF SPONTANEOUS CONTRACTIONS AND TONUS OF HINDGUT MUSCLE.
CC -1- TISSUE SPECIFICITY: BRAIN.
CC -1- MASS SPECTROMETRY: MW=1023.0; METHOD=MALDI.
CC -1- SIMILARITY: SOME SIMILARITY TO TACHYKININS.
KW Tachykinin; Neuropeptide; Amidation.
FT MOD_RES 10
SQ SEQUENCE 10 AA; 1024 MW; C4469D79C9C87DDD CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 6 FF 7

RESULT 32
TRP7_LEUMA
ID TRP7_LEUMA STANDARD; PRT; 10 AA.
AC P81739;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Tachykinin-related peptide 7 (LemTRP 7).
OS Leucophaea maderae (Madeira cockroach).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Orthopteroidea; Dictyoptera; Blattaria;
OC Blaberoidea; Blaberidae; Leucophaea.
RN NCBI_TaxID=6988;
[1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=Brain;
RX MEDLINE=97262666; PubMed=9114447;
RA Muren J.E., Naessel D.R.;
RT "Seven tachykinin-related peptides isolated from the brain of the
RT Madeira cockroach; evidence for tissue-specific expression of
RT isoforms."
RL Peptides 18:7-15(1997).
CC -1- FUNCTION: MYOACTIVE PEPTIDE. INCREASES THE AMPLITUDE AND FREQUENCY
CC OF SPONTANEOUS CONTRACTIONS AND TONUS OF HINDGUT MUSCLE.
CC -1- TISSUE SPECIFICITY: BRAIN.
CC -1- MASS SPECTROMETRY: MW=1023.0; METHOD=MALDI.
CC -1- SIMILARITY: SOME SIMILARITY TO TACHYKININS.
KW Tachykinin; Neuropeptide; Amidation.
FT MOD_RES 10
SQ SEQUENCE 10 AA; 1024 MW; C4469D79C9C87DDD CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 6 FF 7

RESULT 33
TRP7_LEUMA
ID TRP7_LEUMA STANDARD; PRT; 10 AA.
AC P81739;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Tachykinin-related peptide 7 (LemTRP 7).
OS Leucophaea maderae (Madeira cockroach).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Orthopteroidea; Dictyoptera; Blattaria;
OC Blaberoidea; Blaberidae; Leucophaea.
RN NCBI_TaxID=6988;
[1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=Brain;
RX MEDLINE=97262666; PubMed=9114447;
RA Muren J.E., Naessel D.R.;
RT "Seven tachykinin-related peptides isolated from the brain of the
RT Madeira cockroach; evidence for tissue-specific expression of
RT isoforms."
RL Peptides 18:7-15(1997).
CC -1- FUNCTION: MYOACTIVE PEPTIDE. INCREASES THE AMPLITUDE AND FREQUENCY
CC OF SPONTANEOUS CONTRACTIONS AND TONUS OF HINDGUT MUSCLE.
CC -1- TISSUE SPECIFICITY: BRAIN.
CC -1- MASS SPECTROMETRY: MW=1023.0; METHOD=MALDI.
CC -1- SIMILARITY: SOME SIMILARITY TO TACHYKININS.
KW Tachykinin; Neuropeptide; Amidation.
FT MOD_RES 10
SQ SEQUENCE 10 AA; 1024 MW; C4469D79C9C87DDD CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 6 FF 7

RESULT 34
HY7_PIG
ID HY7_PIG STANDARD; PRT; 7 AA.
AC P01153;
DT 21-JUL-1986 (Rel. 01, Created)
DT 21-JUL-1986 (Rel. 01, Last sequence update)
DT 21-JUL-1986 (Rel. 01, Last annotation update)
DE Hypothalamic heptapeptide.
OS Sus scrofa (Pig).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
RN NCBI_TaxID=9823;
[1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=Brain;
RX MEDLINE=97262666; PubMed=9114447;
RA Muren J.E., Naessel D.R.;
RT "Seven tachykinin-related peptides isolated from the brain of the
RT Madeira cockroach; evidence for tissue-specific expression of
RT isoforms."
RL Peptides 18:7-15(1997).
CC -1- FUNCTION: MYOACTIVE PEPTIDE. INCREASES THE AMPLITUDE AND FREQUENCY
CC OF SPONTANEOUS CONTRACTIONS AND TONUS OF HINDGUT MUSCLE.
CC -1- TISSUE SPECIFICITY: BRAIN.
CC -1- MASS SPECTROMETRY: MW=1069.7; METHOD=MALDI.
CC -1- SIMILARITY: SOME SIMILARITY TO TACHYKININS.
KW Tachykinin; Neuropeptide; Amidation.
FT MOD_RES 10
SQ SEQUENCE 10 AA; 1068 MW; C4541679C9C865BD CRC64;

Query Match 21.8%; Score 12; DB 1; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.1e+04;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FF 8
Db 6 FF 7

RESULT 35
CHOX_ALCSP
ID CHOX_ALCSP STANDARD; PRT; 7 AA.
AC P16101;
DT 01-APR-1990 (Rel. 14, Created)
DT 01-APR-1990 (Rel. 14, Last sequence update)
DT 01-APR-1990 (Rel. 14, Last annotation update)
DE Choline oxidase (EC 1.1.3.17) (Fragment).
OS Alcaligenes sp.
OC Bacteria; Proteobacteria; beta subdivision; Alcaligenaceae;
OC Alcaligenes.
OX NCBI_TaxID=512;
[1]
RP SEQUENCE.
RX MEDLINE=81006769; PubMed=6997283;
RA Ohka-Fukuyama M., Miyake Y., Emi S., Yamano T.;
RT "Identification and properties of the prosthetic group of choline
RT oxidase from Alcaligenes sp."
RL J. Biochem. 88:197-203(1980).
CC -1- CATALYTIC ACTIVITY: Choline + O(2) = betaine aldehyde + H(2)O(2).
KW PIR: A15398; A15398.
DR Oxidoreductase.
FT NON_TER 7
SQ SEQUENCE 7 AA; 839 MW; 7415B1E457644AC0 CRC64;

Query Match 20.0%; Score 11; DB 1; Length 7;
Best Local Similarity 25.0%; Pred. No. 1e+05;
Matches 1; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

Qy 1 HHQK 4
Db 4 NHSR 7
```

RN SEQUENCE, AND SYNTHESIS.
 RX MEDLINE=81213980; PubMed=6263778;
 RA Chang R.C.C., Huang W.-Y., Aizimura A., Redding T.W., Coy D.H.,
 RA Saffran M., Kong A., Hamilton J.W., Conn D.V., Schally A.V.;
 RT "Isolation, structure and synthesis of a heptapeptide with in vitro
 RT ACTH-releasing activity from porcine hypothalamus.";
 RL Horm. Metab. Res. 13:228-232(1981).
 DR PIR: A01417; NYPG7.
 SQ SEQUENCE 7 AA; 957 MW; 632B45B1FB5059A0 CRC64;

 Query Match 20.0%; Score 11; DB 1; Length 7;
 Best Local Similarity 50.0%; Pred. No. 1e+05;
 Matches 2; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

 QY 1 HHQK 4
 DB 4 HSYK 7

 RESULT 35
 UF03_MOUSE STANDARD; PRT; 7 AA.
 AC P38641;
 DT 01-OCT-1994 (Rel. 30, Created)
 DT 01-OCT-1994 (Rel. 30, Last sequence update)
 DT 01-FEB-1995 (Rel. 31, Last annotation update)
 DE Unknown protein from 2D-page of fibroblasts (P36) (Fragment).
 OS Mus musculus (Mouse).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 OX NCBI_TaxID=10090;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Fibroblast; PubMed=7523108;
 RX MEDLINE=95009907; PubMed=7523108;
 RA Merrick B.A., Patterson R.M., Wichter L.L., He C., Salkirk J.K.;
 RT "Separation and sequencing of familial and novel murine proteins
 RT using preparative two-dimensional gel electrophoresis.";
 RL Electrophoresis 15:735-745(1994).
 CC -1- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
 CC PROTEIN IS: 5.1, ITS MW IS: 36 kDa.
 FT NON_TER 7
 SQ SEQUENCE 7 AA; 842 MW; 6AA72B1DBD1B1180 CRC64;

 Query Match 20.0%; Score 11; DB 1; Length 7;
 Best Local Similarity 33.3%; Pred. No. 1e+05;
 Matches 1; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

 QY 2 HQK 4
 DB 1 HEE 3

 RESULT 36
 AKH_TABAT STANDARD; PRT; 8 AA.
 AC P14595;
 DT 01-JAN-1990 (Rel. 13, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Adipokinetic hormone (AKH) (Dipteran corpora cardiaca factor I)
 DE (DCC 1).
 OS Tabanus atratus (Horse fly).
 OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
 OC Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Tabanomorpha;
 OC Tabanidae; Tabanus.
 OX NCBI_TaxID=7207;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Corpora cardiaca;
 RX MEDLINE=90046758; PubMed=2813385;
 RA Jaffe H., Raina A.K., Riley C.T., Fraser B.A., Nachman R.J.,

RA Vogel V.W., Zhang Y.-S., Hayes D.K.;
 RT "Primary structure of two neuropeptide hormones with adipokinetic and
 RT hypotrehalosemic activity isolated from the corpora cardiaca of horse
 RT flies (Diptera).";
 RL Proc. Natl. Acad. Sci. U.S.A. 86:8161-8164(1989).
 CC -1- FUNCTION: THIS HORMONE, RELEASED FROM CELLS IN THE CORPORA
 CC CARDIACA AFTER THE BEGINNING OF FLIGHT, CAUSES RELEASE OF
 CC DIGLYCERIDES FROM THE FAT BODY AND THEN STIMULATES THE FLIGHT
 CC MUSCLES TO USE THESE DIGLYCERIDES AS AN ENERGY SOURCE.
 CC -1- SIMILARITY: BELONGS TO THE AKH / HETH / RPCH FAMILY.
 DR PIR: A33995; A33995.
 DR InterPro: IPR002047; AKH.
 DR PROSITE: PS00256; AKH; 1.
 KW Neuropeptide; Amidation; Flight.
 FT MOD_RES 1 1 PYRROLIDONE CARBOXYLIC ACID.
 FT MOD_RES 8 8 AMIDATION.
 SQ SEQUENCE 8 AA; 949 MW; 86786771A9D1A736 CRC64;

 Query Match 20.0%; Score 11; DB 1; Length 8;
 Best Local Similarity 50.0%; Pred. No. 1e+05;
 Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

 QY 4 KLVF 7
 DB 1 QLTF 4

 RESULT 37
 HTF2_PERAM STANDARD; PRT; 8 AA.
 AC P04549;
 DT 13-AUG-1987 (Rel. 05, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Hypertrahalosemic factor II (Neuropeptide M-II) (Periplanetin CC-2)
 DE (PeA-CAH-II) (Led-CC-II) (Hypertrahalosemic neuropeptide II).
 OS Periplaneta americana (American cockroach).
 OS Leptinotarsa decemlineata (Colorado potato beetle), and
 OS Blattella orientalis (Oriental cockroach).
 OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
 OC Pterygota; Neoptera; Orthopteroidea; Dictyoptera; Blattaria;
 OC Blattoidea; Blattidae; Periplaneta.
 OX NCBI_TaxID=6978, 7539, 6976;
 RN [1]
 RP SEQUENCE.
 RC SPECIES=P.americana;
 RX MEDLINE=85046530; PubMed=6548628;
 RA Witten J.L., Schaffer M.H., O'Shea M., Cook J.C., Hemling M.E.,
 RA Rinehart K.L., Jr.;
 RT "Structures of two cockroach neuropeptides assigned by fast atom
 RT bombardment mass spectrometry.";
 RL Biochem. Biophys. Res. Commun. 124:350-358(1984).
 RN [2]
 RP SEQUENCE.
 RC SPECIES=P.americana;
 RX MEDLINE=84298179; PubMed=6591205;
 RA Scarborough R.M., Jamieson G.C., Kalish F., Kramer S.J., McEnroe G.A.,
 RA Miller C.A., Schooley D.A.;
 RT "Isolation and primary structure of two peptides with
 RT cardioacceleratory and hyperglycemic activity from the corpora
 RT cardiaca of Periplaneta americana.";
 RL Proc. Natl. Acad. Sci. U.S.A. 81:5575-5579(1984).
 RN [3]
 RP SEQUENCE.
 RC SPECIES=L.decemlineata; TISSUE=Corpora cardiaca;
 RX MEDLINE=90160053; PubMed=2576128;
 RA Gaede G., Kellner R.;
 RT "The metabolic neuropeptides of the corpus cardiaca from the potato
 RT beetle and the American cockroach are identical.";
 RL Peptides 10:1287-1289(1989).
 RN [4]
 RP SEQUENCE.
 RC SPECIES=B.orientalis; TISSUE=Corpora cardiaca;

RX MEDLINE=90253659; PubMed=2340112;
 RA Gaede G., Rinehart K.L. Jr.;
 RT "Primary structures of hypertrehalosemic neuropeptides isolated from
 the corpora cardiaca of the cockroaches *Leucophaea maderae*,
Gromphadorhina portentosa, *Blattella germanica* and *Blatta orientalis*
 and of the stick insect *Extatosoma tiaratum* assigned by tandem fast
 atom bombardment mass spectrometry.";
 RL Biol. Chem. Hoppe-Seyler 371:345-354(1990).
 CC -!- FUNCTION: HYPERTREHALOSEMIC FACTORS ARE NEUROPEPTIDES THAT
 ELEVATE THE LEVEL OF TREHALOSE IN THE HEMOLYMPH OF INSECTS).
 CC THE MAJOR CARBOHYDRATE IN THE HEMOLYMPH OF INSECTS).
 CC -!- SIMILARITY: BELONGS TO THE AKH / HRTH / RPCH FAMILY.
 DR PIR; A05170;
 DR PIR; A05170;
 DR PIR; S08996; S08996.
 DR PIR; B44960; B44960.
 DR PIR; B49823; B49823.
 DR InterPro; IPR002047; AKH.
 DR PROSITE; PS00256; AKH; 1.
 KW Neuropeptide; Amidation.
 FT MOD_RES 1 1
 FT MOD_RES 8 8 PYRROLIDONE CARBOXYLIC ACID.
 SQ SEQUENCE 8 AA; 1006 MW; 86745771A9DIA736 CRC64;
 Amidation.
 Query Match 20.0%; Score 11; DB 1; Length 8;
 Best Local Similarity 50.0%; Pred. No. 1e+05;
 Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;
 QY 4 KLVF 7
 Db :|||
 1 QLTFF 4
 RESULT 38
 FAR6_PANRE
 ID FAR6_PANRE STANDARD; PRT; 9 AA.
 AC P82661;
 DT 16-OCT-2001 (Rel. 40, Created)
 DT 16-OCT-2001 (Rel. 40, Last sequence update)
 DE FMRFamide-like neuropeptide PF5 (AMRNALVRF-amide).
 OS Panagrellus redivivus.
 OC Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida;
 OC Panagrolaimoidea; Panagrolaimidae; Panagrellus.
 OX NCBI_TaxID=6233;
 RN [1]
 RP SEQUENCE, FUNCTION, AND AMIDATION.
 RA Moffet C.L., Marks N.J., Halton D.W., Thomson D.P., Geary T.G.,
 RA Maule A.G.;
 RT "Isolation, characterization and pharmacology of FMRFamide-related
 peptides (FARPs) from free-living nematode, *Panagrellus redivivus*,";
 RL Submitted (JUL-2000) to the SWISS-PROT data bank.
 CC -!- FUNCTION: MYOACTIVE.
 CC -!- SIMILARITY: BELONGS TO THE FARP (FMRFAMIDE RELATED PEPTIDE)
 FAMILY.
 KW Neuropeptide; Amidation.
 FT MOD_RES 9 9
 FT MOD_RES 9 9 AMIDATION.
 SQ SEQUENCE 9 AA; 1077 MW; A0D112C72DD45406 CRC64;
 Amidation.
 Query Match 20.0%; Score 11; DB 1; Length 9;
 Best Local Similarity 75.0%; Pred. No. 1e+05;
 Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;
 QY 5 LVFF 8
 Db :|||
 6 LVRF 9
 RESULT 39
 ULAK_MOUSE
 ID ULAK_MOUSE STANDARD; PRT; 9 AA.
 AC P99031;
 DT 15-DEC-1998 (Rel. 37, Created)
 DT 15-DEC-1998 (Rel. 37, Last sequence update)

DT 15-DEC-1998 (Rel. 37, Last annotation update)
 DE Unknown protein from 2D-page of liver tissue (Spot 2D-0014LD)
 DE (Fragment).
 OS *Mus musculus* (Mouse).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 OX NCBI_TaxID=10090;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Liver;
 RA Sanchez J.-C., Rouge V., Frutiger S., Hughes G.J., Yan J.X.,
 RA Hoogland C., Appel R.D., Binz P.-A., Hochstrasser D.F.,
 RA Cowthorne M.;
 RL Submitted (AUG-1998) to the SWISS-PROT data bank.
 CC -!- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
 PROTEIN IS: 6.0, ITS MW IS: 12.5 kDa.
 CC SWISS-2DPAGE; P99031; MOUSE.
 DR NON_TER 9
 SQ SEQUENCE 9 AA; 1106 MW; E1E842C3240B145A CRC64;
 Amidation.
 Query Match 20.0%; Score 11; DB 1; Length 9;
 Best Local Similarity 16.7%; Pred. No. 1e+05;
 Matches 1; Conservative 4; Mismatches 1; Indels 0; Gaps 0;
 QY 1 HHQKV 6
 Db :|||
 3 NERKVI 8
 RESULT 40
 BPP2_BOTIN
 ID BPP2_BOTIN STANDARD; PRT; 10 AA.
 AC P30422;
 DT 01-APR-1993 (Rel. 25, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Bradykinin-potentiating peptide S4,3,1 (10C) (Angiotensin-converting
 enzyme inhibitor).
 OS Bothrops insularis (Island jararaca) (Queimada jararaca).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Lepidosauria; Squamata; Scieroglossa; Serpentes; Colubroidea;
 OC Viperidae; Crotalinae; Bothrops.
 OX NCBI_TaxID=8723;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Venom;
 RX MEDLINE=90351557; PubMed=2386615;
 RA Cintra A.C.O., Vieira C.A., Giglio J.R.;
 RT "Primary structure and biological activity of bradykinin potentiating
 peptides from *Bothrops insularis* snake venom.";
 RL J. Protein Chem. 9:221-227(1990).
 CC -!- FUNCTION: THIS PEPTIDE BOTH INHIBITS THE ACTIVITY OF THE
 ANGIOTENSIN-CONVERTING ENZYME AND ENHANCES THE ACTION OF
 BRADYKININ BY INHIBITING THE KINASES THAT INACTIVATE IT.
 CC IT ACTS AS AN INDIRECT HYPOTENSIVE AGENT.
 DR PIR; B37196; B37196.
 KW Hypotensive agent; Venom.
 FT MOD_RES 1 1
 FT MOD_RES 1 1 PYRROLIDONE CARBOXYLIC ACID.
 SQ SEQUENCE 10 AA; 1213 MW; 30C53546C761F773 CRC64;
 Amidation.
 Query Match 20.0%; Score 11; DB 1; Length 10;
 Best Local Similarity 66.7%; Pred. No. 1.7e+04;
 Matches 2; Conservative 0; Mismatches 1; Indels 0; Gaps 0;
 QY 1 HHQ 3
 Db :|||
 5 HFQ 7
 RESULT 41
 FAR6_PANRE
 ID FAR6_PANRE STANDARD; PRT; 10 AA.
 AC P82660;

DT 16-OCT-2001 (Rel. 40, Created)
 DT 16-OCT-2001 (Rel. 40, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE FMRamide-like neuropeptide PF6 (NGAPQPFVRF-amide).
 OS Panagrellus redivivus.
 OC Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida;
 OC Panagrolaimidae; Panagrolaimidae; Panagrellus.
 OX NCBI_TaxID=6233;
 [1]
 RP SEQUENCE, FUNCTION, AND AMIDATION.
 RA Moffett C.L., Marks N.J., Halton D.W., Thomson D.P., Geary T.G.,
 RA Maule A.G.;
 RT "Isolation, characterization and pharmacology of RMRamide-related
 RT peptides (FARPs) from free-living nematode, Panagrellus redivivus";
 RL Submitted (JUL-2000) to the SWISS-PROT data bank.
 CC -1- FUNCTION: MYOACTIVE.
 CC -1- SIMILARITY: BELONGS TO THE FARP (FMRAMIDE RELATED PEPTIDE)
 CC FAMILY.
 KW Neuropeptide; Amidation.
 FT MOD_RES 10 10 AMIDATION.
 SQ SEQUENCE 10 AA; 1132 MW; CB13E4C9D776C76D CRC64;

 Query Match 20.08; Score 11; DB 1; Length 10;
 Best Local Similarity 50.08; Pred. No. 1.7e+04;
 Matches 3; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

 QY 3 QKLVFF 8
 DB 5 QPFVRF 10

 RESULT 42
 GONL_PETMA STANDARD; PRT; 10 AA.
 AC P04378;
 DT 20-MAR-1987 (Rel. 04, Created)
 DT 20-MAR-1987 (Rel. 04, Last sequence update)
 DT 01-JUN-1994 (Rel. 29, Last annotation update)
 DE Gonadoliberin I (Gonadotropin-releasing hormone I) (GNRH-I)
 DE (Luliberin I).
 OS Petromyzon marinus (Sea lamprey).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Hyperoartia;
 OC Petromyzontiformes; Petromyzontidae; Petromyzon.
 OX NCBI_TaxID=7757;
 [1]
 RP SEQUENCE.
 RC TISSUE=Brain;
 RA Sherwood N.M., Sower S.A., Marshak D.R., Fraser B.A., Brownstein M.J.;
 RT "Primary structure of gonadotropin-releasing hormone from lamprey
 RT brain.";
 RL J. Biol. Chem. 261:4812-4819(1986).
 CC -1- FUNCTION: STIMULATES THE SECRETION OF BOTH LUTEINIZING AND
 CC FOLLICLE-STIMULATING HORMONES.
 CC -1- SUBCELLULAR LOCATION: Secreted.
 CC -1- SIMILARITY: BELONGS TO THE GNRH FAMILY.
 PIR: A01412; RHLMGs.
 DR InterPro: IPR002012; GNRH.
 DR Pfam: PF00446; GNRH; 1.
 DR PROSITE; PS00473; GNRH; 1.
 KW Hormone; Amidation; Hypothalamus.
 FT MOD_RES 1 1 PYROLIDONE CARBOXYLIC ACID.
 FT MOD_RES 10 10 AMIDATION.
 SQ SEQUENCE 10 AA; 1244 MW; 1E4B36237B1735AB CRC64;

 Query Match 20.08; Score 11; DB 1; Length 10;
 Best Local Similarity 50.08; Pred. No. 1.7e+04;
 Matches 2; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

 QY 2 HQKL 5
 DB 2 HYSL 5

RESULT 43
 HTF2_CARMO STANDARD; PRT; 10 AA.
 AC P11385;
 DT 01-JUL-1989 (Rel. 11, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 30-MAY-2000 (Rel. 39, Last annotation update)
 DE Hypertrehalosaemic factor II (HTF-II) (HRTH-II) (Hypertrehalosaemic
 DE neuropeptide II).
 OS Carausius morosus (Indian stick insect), and
 OS Extatosoma tiaratum (Stick insect).
 OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
 OC Plerygota; Neoptera; Orthopteroidea; Phasmatodea; Heteronemiidae;
 OC Carausius.
 OX NCBI_TaxID=7022, 7024;
 [1]
 RP SEQUENCE.
 RC SPECIES=C.morosus; TISSUE=Corpora cardiaca;
 RX MEDLINE=87157103; PubMed=3828078;
 RA Gaede G., Rinehart K.L. Jr.;
 RT "Primary structure of the hypertrehalosaemic factor II from the
 RT corpus cardiacum of the Indian stick insect, Carausius morosus,
 RT determined by fast atom bombardment mass spectrometry.";
 RL Biol. Chem. Hoppe-Seyler 368:67-75(1987).
 RN [2]
 RP SEQUENCE.
 RC SPECIES=E.tiaratum; TISSUE=Corpora cardiaca;
 RX MEDLINE=90253659; PubMed=2340112;
 RA Gaede G., Rinehart K.L. Jr.;
 RT "Primary structures of hypertrehalosaemic neuropeptides isolated from
 RT the corpora cardiaca of the cockroaches Leucophaea maderae,
 RT Gromphadorhina portentosa, Blattella germanica and Blattella orientalis
 RT and of the stick insect Extatosoma tiaratum assigned by tandem fast
 RT atom bombardment mass spectrometry.";
 RL Biol. Chem. Hoppe-Seyler 371:345-354(1990).
 RN [3]
 RP CARBOHYDRATE-LINKAGE SITE.
 RC SPECIES=C.morosus; TISSUE=Corpora cardiaca;
 RX MEDLINE=93129188; PubMed=1482345;
 RA Gaede G., Kellner R., Rinehart K.L. Jr., Proefke M.L.;
 RT "A tryptophan-substituted member of the AKH/RPCH family isolated from
 RT a stick insect corpus cardiacum.";
 RL Biochem. Biophys. Res. Commun. 189:1303-1309(1992).
 CC -1- FUNCTION: HYPERTREHALOSAEMIC FACTORS ARE NEUROPEPTIDES THAT
 CC ELEVATE THE LEVEL OF TREHALOSE IN THE HEMOLymph OF INSECTS).
 CC THE MAJOR CARBOHYDRATE IN THE HEMOLymph OF INSECTS).
 CC -1- MASS SPECTROMETRY: MW=1308.61; METHOD=FAb.
 CC -1- SIMILARITY: BELONGS TO THE AKH / HRTH / RPCH FAMILY.
 PIR: S07157; S07157.
 DR InterPro: IPR002047; AKH.
 DR PROSITE; PS00256; AKH; 1.
 KW Neuropeptide; Amidation; Glycoprotein.
 FT MOD_RES 1 1 PYROLIDONE CARBOXYLIC ACID.
 FT MOD_RES 8 8 C-LINKED (MAN) (PROBABLE).
 FT MOD_RES 10 10 AMIDATION.
 SQ SEQUENCE 10 AA; 1164 MW; 9B9036745771A9D1 CRC64;

 Query Match 20.08; Score 11; DB 1; Length 10;
 Best Local Similarity 50.08; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

 QY 4 KLVF 7
 DB 1 QLTF 4

 RESULT 44
 HTF_HELZE STANDARD; PRT; 10 AA.
 ID HTF_HELZE
 AC P16353;
 DT 01-AUG-1990 (Rel. 15, Created)

DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Hypertrehalosemic hormone (He2-HRTH).
 OS Heliothis zea (Corn earworm) (Bollworm).
 OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
 OC Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia;
 OC Noctuoidea; Noctuidae; Heliothinae; Helicoverpa.
 OX NCBI_TaxID=7113;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Corpora cardiaca;
 RX MEDLINE=86326324; PubMed=3415690;
 RA Jaffe H., Raina A.K., Riley C.T., Fraser B.A., Bird T.G.,
 RA Tseng C.M., Zhang Y.S., Hayes D.K.;
 RT "Isolation and primary structure of a neuropeptide hormone from
 RT Heliothis zea with hypertrehalosemic and adipokinetic activities.";
 RL Biochem. Biophys. Res. Commun. 155:344-350(1988).
 CC -!- FUNCTION: HYPERTREHALOSEMIC FACTORS ARE NEUROPEPTIDES THAT
 CC ELEVATE THE LEVEL OF TREHALOSE IN THE HEMOLYMPH (TREHALOSE IS
 CC THE MAJOR CARBOHYDRATE IN THE HEMOLYMPH OF INSECTS).
 CC -!- SIMILARITY: BELONGS TO THE AKH / HRTH / RPCH FAMILY.
 CC PIR: A31571; A31571.
 DR InterPro: IPR002047; AKH.
 DR PROSITE: PS00256; AKH; 1.
 KW Neuropeptide; Amidation.
 FT MOD_RES 1 1
 FT MOD_RES 10 10 PYRROLIDONE CARBOXYLIC ACID.
 FT SEQUENCE 10 AA; 1096 MW; 8E70367865A5B9D1 CRC64;
 SQ SEQUENCE 10 AA; 1096 MW; 8E70367865A5B9D1 CRC64;
 Query Match 20.0%; Score 11; DB 1; Length 10;
 Best Local Similarity 50.0%; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;
 QY 4 KLVFF 7
 Db :| |
 1 QLTFF 4

RESULT 45
 HTF_TABAT
 ID HTF_TABAT STANDARD; PRT; 10 AA.
 AC P14596;
 DT 01-JAN-1990 (Rel. 13, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Hypertrehalosemic factor (HOTH) (Dipteran corpora cardiaca factor II)
 DE (DCC II).
 OS Tabanus atratus (Horse fly).
 OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
 OC Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Tabanomorpha;
 OC Tabanidae; Tabanus.
 OX NCBI_TaxID=7207;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Corpora cardiaca;
 RX MEDLINE=90046758; PubMed=2813385;
 RA Jaffe H., Raina A.K., Riley C.T., Fraser B.A., Nachman R.J.,
 RA Vogel V.W., Zhang Y.-S., Hayes D.K.;
 RT "Primary structure of two neuropeptide hormones with adipokinetic and
 RT hypotrehalosemic activity isolated from the corpora cardiaca of horse
 RT flies (Diptera).";
 RL Proc. Natl. Acad. Sci. U.S.A. 86:8161-8164(1989).
 CC -!- FUNCTION: HYPERTREHALOSEMIC FACTORS ARE NEUROPEPTIDES THAT
 CC ELEVATE THE LEVEL OF TREHALOSE IN THE HEMOLYMPH (TREHALOSE IS
 CC THE MAJOR CARBOHYDRATE IN THE HEMOLYMPH OF INSECTS).
 CC -!- SIMILARITY: BELONGS TO THE AKH / HRTH / RPCH FAMILY.
 CC PIR: B33995; B33995.
 DR InterPro: IPR002047; AKH.
 DR PROSITE: PS00256; AKH; 1.
 KW Neuropeptide; Amidation.
 FT MOD_RES 1 1
 FT MOD_RES 10 10 PYRROLIDONE CARBOXYLIC ACID.
 FT SEQUENCE 10 AA; 1169 MW; 916036786771A9D1 CRC64;
 SQ SEQUENCE 10 AA; 1169 MW; 916036786771A9D1 CRC64;

Query Match 20.0%; Score 11; DB 1; Length 10;
 Best Local Similarity 50.0%; Pred. No. 1.7e+04;
 Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;
 QY 4 KLVFF 7
 Db :| |
 1 QLTFF 4

RESULT 46
 Q2OB_COMTE
 ID Q2OB_COMTE STANDARD; PRT; 10 AA.
 AC P80465;
 DT 01-NOV-1995 (Rel. 32, Created)
 DT 01-NOV-1995 (Rel. 32, Last sequence update)
 DT 16-OCT-2001 (Rel. 40, Last annotation update)
 DE Quinoline 2-oxidoreductase, beta chain [EC 1.3.99.17] (Fragment).
 OS Comamonas testosteroni (Pseudomonas testosteroni).
 OC Bacteria; Proteobacteria; beta subdivision; Comamonadaceae; Comamonas.
 OX NCBI_TaxID=285;
 RN [1]
 RP SEQUENCE.
 RC STRAIN=63;
 RX MEDLINE=96035889; PubMed=7556204;
 RA Schach S., Tshisuaka B., Fetzner S., Lingens F.;
 RT "Quinoline 2-oxidoreductase and 2-oxo-1,2-dihydroquinoline 5,6-
 RT dioxygenase from Comamonas testosteroni 63. The first two enzymes in
 RT quinoline and 3-methylquinoline degradation.";
 RL Eur. J. Biochem. 232:536-544(1995).
 CC -!- FUNCTION: CONVERTS (3-METHYL-)-QUINOLINE TO (3-METHYL-)-2-OXO-
 CC 1,2-DIHYDROQUINOLINE.
 CC -!- CATALYTIC ACTIVITY: Quinoline + acceptor + H(2)O = isoquinolin-
 CC 1(2H)-one + reduced acceptor.
 CC -!- COFACTOR: FAD, MOLYBDENUM AND IRON-SULFUR.
 CC -!- PATHWAY: FIRST STEP IN THE DEGRADATION OF QUINOLINE AND
 CC (3-METHYL-)-QUINOLINE.
 CC -!- SUBUNIT: HETEROHEXAMER OF TWO ALPHA CHAINS, TWO BETA CHAINS, AND
 CC TWO GAMMA CHAINS (PROBABLE).
 KW Oxidoreductase; Flavoprotein; FAD; Molybdenum.
 FT NON_TER 10 10
 FT SEQUENCE 10 AA; 1241 MW; C2E2C25DD9CDC769 CRC64;
 SQ SEQUENCE 10 AA; 1241 MW; C2E2C25DD9CDC769 CRC64;
 Query Match 20.0%; Score 11; DB 1; Length 10;
 Best Local Similarity 50.0%; Pred. No. 1.7e+04;
 Matches 3; Conservative 0; Mismatches 3; Indels 0; Gaps 0;
 QY 4 KLVFFA 9
 Db | | |
 2 KFPFA 7

RESULT 47
 BPP7_BOTIN
 ID BPP7_BOTIN STANDARD; PRT; 5 AA.
 AC P30425;
 DT 01-APR-1993 (Rel. 25, Created)
 DT 01-FEB-1994 (Rel. 28, Last sequence update)
 DT 01-FEB-1994 (Rel. 28, Last annotation update)
 DE Bradykinin-potentiating peptide S5,2 (5A) (Angiotensin-converting
 DE enzyme inhibitor).
 OS Bothrops insularis (Island jararaca) (Queimada jararaca).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Lepidodactylia; Squamata; Serpentes; Colubroidea;
 OC Viperidae; Crotalinae; Bothrops.
 OX NCBI_TaxID=8723;
 RN [1]
 RP SEQUENCE.
 RC TISSUE=Venom;
 RX MEDLINE=90351557; PubMed=2386615;
 RA Cintra A.C.O., Vieira C.A., Giglio J.R.;
 RT "Primary structure and biological activity of bradykinin potentiating
 RT peptides from Bothrops insularis snake venom.";

RL J. Protein Chem. 9:221-227(1990).
CC -!- FUNCTION: THIS PEPTIDE BOTH INHIBITS THE ACTIVITY OF THE
CC ANGIOTENSIN-CONVERTING ENZYME AND ENHANCES THE ACTION OF
CC BRADYKININ BY INHIBITING THE KINASES THAT INACTIVATE IT.
CC IT ACTS AS AN INDIRECT HYPOTENSIVE AGENT.
DR PIR: G37196; G37196.
KW Hypotensive agent; Venom.
FT MOD_RES 1 1
SQ SEQUENCE 5 AA; 629 MW; 776DC37326B00000 CRC64;

Query Match 18.2%; Score 10; DB 1; Length 5;
Best Local Similarity 100.0%; Pred. No. 1e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 3 QK 4
| |
Db 1 QK 2

RESULT 48

ACI_THUAL STANDARD; PRT; 8 AA.
AC P16691;
DT 01-NOV-1990 (Rel. 16, Created)
DT 01-NOV-1990 (Rel. 16, Last sequence update)
DT 01-NOV-1990 (Rel. 16, Last annotation update)
DE Angiotensin-converting enzyme inhibitor.
OS Thunus albacares (Yellowfin tuna) (Neothunnus macropterus).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
OC Acanthomorpha; Acanthopterygii; Percomorpha; Perciformes; Scombroidei;
OC Scombridae; Thunnus.
OX NCBI_TaxID=8236;
RN [1]
RP SEQUENCE.

RC TISSUE=Muscle;
RX MEDLINE=88326322; PubMed=3415688;
RA Kohama Y., Matsumoto S., Oka H., Teramoto T., Okabe M., Mimura T.;
RT "Isolation of angiotensin-converting enzyme inhibitor from tuna muscle."
RL Biochem. Biophys. Res. Commun. 155:332-337(1988).
DR PIR: A31570; A31570.
SQ SEQUENCE 8 AA; 953 MW; 6AA863733051F1B7 CRC64;

Query Match 18.2%; Score 10; DB 1; Length 8;
Best Local Similarity 66.7%; Pred. No. 1e+05;
Matches 2; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

Qy 2 HOK 4
| |
Db 3 HIK 5

RESULT 49

ALLI_CYPDPO STANDARD; PRT; 8 AA.
AC P82152;
DT 30-MAY-2000 (Rel. 39, Created)
DT 30-MAY-2000 (Rel. 39, Last sequence update)
DT 30-MAY-2000 (Rel. 39, Last annotation update)
DE Cydia statin 1.
OS Cydia pomonella (Codling moth).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia;
OC Tortricidae; Tortricidae; Olethreutinae; Cydia.
OX NCBI_TaxID=82600;
RN [1]
RP SEQUENCE.

RC TISSUE=Larva;
RX MEDLINE=98054539; PubMed=9392829;
RA Duve H., Johnsen A.H., Maestro J.-L., Scott A.G., Winstanley D.,
RA Davey M., East P.D., Thorpe A.;
RT "Lepidopteran peptides of the allatostatin superfamily."

RL Peptides 18:1301-1309(1997).
CC -!- SIMILARITY: BELONGS TO THE ALLATOSTATIN FAMILY.
CC NEUROPEPTIDE; Amidation.
FT MOD_RES 8
SQ SEQUENCE 8 AA; 934 MW; C82879C45B51F775 CRC64;
Query Match 18.2%; Score 10; DB 1; Length 8;
Best Local Similarity 50.0%; Pred. No. 1e+05;
Matches 1; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HH 2
| |
Db 3 HY 4

RESULT 50

LCK8_LEUMA STANDARD; PRT; 8 AA.
AC P19990;
DT 01-FEB-1991 (Rel. 17, Created)
DT 01-FEB-1991 (Rel. 17, Last sequence update)
DT 01-FEB-1991 (Rel. 17, Last annotation update)
DE Leucokinin VIII (L-VIII).
OS Leucophaea maderae (Madeira cockroach).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Orthopteroidea; Dictyoptera; Blattaria;
OC Blaberoidea; Blaberidae; Leucophaea.
OX NCBI_TaxID=6988;
RN [1]
RP SEQUENCE.
RC TISSUE=Head;
RA Holman G.M., Cook B.J., Nachman R.J.;
RT "Isolation, primary structure and synthesis of leucokinin VII and VIII: the final members of this new family of cephalomyotropic peptides isolated from head extracts of Leucophaea maderae."
RL Comp. Biochem. Physiol. 88C:31-34(1987).
CC -!- FUNCTION: THIS CEPHALOMYOTROPIC PEPTIDE STIMULATES CONTRACTILE
CC ACTIVITY OF COCKROACH PROTODEUM (HINDGUT).
CC -!- SIMILARITY: TO THE OTHER LEUCOKININS.
DR PIR: JS0318; JS0318.
KW Neuropeptide; Amidation.
FT MOD_RES 8
SQ SEQUENCE 8 AA; 902 MW; 736365AB59CAADD8 CRC64;

Query Match 18.2%; Score 10; DB 1; Length 8;
Best Local Similarity 33.3%; Pred. No. 1e+05;
Matches 1; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
| |
Db 4 FYS 6

Search completed: October 29, 2002, 09:37:57
Job time : 13 secs

GenCore version 5.1.3
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OM protein - protein search, using sw model
Run on: October 29, 2002, 09:32:02 ; Search time 24 Seconds
(without alignments)
72.081 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLFFFAE 10
Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5
Searched: 562222 seqs, 172994929 residues
Total number of hits satisfying chosen parameters: 1088

Minimum DB seq length: 0
Maximum DB seq length: 10
Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : SPTREMBL_19:*
1: sp_archaea:*
2: sp_bacteria:*
3: sp_fungi:*
4: sp_human:*
5: sp_invertebrate:*
6: sp_mammal:*
7: sp_mhc:*
8: sp_organelle:*
9: sp_phase:*
10: sp_plant:*
11: sp_rodent:*
12: sp_virus:*
13: sp_vertebrate:*
14: sp_unclassified:*
15: sp_rvirus:*
16: sp_bacteriaph:*
17: sp_archaeap:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	SUMMARIES			Description
	Score	Query Match %	ID	
1	21	38.2	9 10 P82440	P82440 nicotiana t
2	19	34.5	9 8 Q9GD36	Q9GD36 juncus effu
3	18	32.7	9 11 Q924N8	Q924N8 mus musculus
4	17	30.9	10 12 Q39952	Q39952 hepatitis g
5	17	30.9	10 12 Q9WLE4	Q9WLE4 hepatitis g
6	16	29.1	5 13 P82070	P82070 litorea rub
7	16	29.1	5 13 P82071	P82071 litorea rub
8	16	29.1	8 4 Q15894	Q15894 homo sapien
9	16	29.1	8 10 Q40530	Q40530 nicotiana t
10	16	29.1	9 2 Q47556	Q47556 escherichia
11	16	29.1	9 4 Q9H4M8	Q9H4M8 homo sapien
12	15	27.3	8 2 Q9R5L7	Q9R5L7 clostridium
13	15	27.3	8 3 Q13591	Q13591 saccharomyc
14	15	27.3	8 4 Q15889	Q15889 homo sapien
15	15	27.3	9 2 Q30790	Q30790 erwinia amy
16	15	27.3	9 2 Q46179	Q46179 clostridium

17	15	27.3	9	6	Q9TT77	Q9tt77 bos taurus
18	15	27.3	9	12	Q92766	Q92766 canine dist
19	15	27.3	10	2	Q9RJ38	Q9rj38 helicobacte
20	15	27.3	10	4	Q9UCS3	Q9ucs3 homo sapien
21	15	27.3	10	5	P82223	P82223 bombyx mori
22	15	27.3	10	5	P82224	P82224 bombyx mori
23	14	25.5	8	4	Q9UMH9	Q9umh9 homo sapien
24	14	25.5	9	4	Q9UC36	Q9uc36 homo sapien
25	14	25.5	9	8	Q94VG2	Q94vg2 varanus ind
26	14	25.5	10	10	P82937	P82937 hordeum vul
27	13	23.6	8	2	O09258	O09258 synechococ
28	13	23.6	8	2	Q9S6D5	Q9s6d5 escherichia
29	13	23.6	8	2	Q56759	Q56759 xanthobacte
30	13	23.6	8	4	Q9UD24	Q9ud24 homo sapien
31	13	23.6	8	5	O15899	O15899 babesia ovi
32	13	23.6	8	6	Q9GMH3	Q9gmh3 lagenorhync
33	13	23.6	8	6	Q28866	Q28866 megaptera n
34	13	23.6	8	6	O02831	O02831 oryctolagus
35	13	23.6	8	11	Q99NX9	Q99nx9 hydrochoeru
36	13	23.6	9	4	O15891	O15891 homo sapien
37	13	23.6	9	6	Q9GJV3	Q9gJV3 lagenorhync
38	13	23.6	9	6	Q9GJV2	Q9gJV2 lagenorhync
39	13	23.6	9	6	Q9GJV1	Q9gJV1 lagenorhync
40	13	23.6	9	8	Q9T688	Q9t688 gecko gecko
41	13	23.6	10	8	Q9T4P9	Q9t4p9 liolaemus d
42	13	23.6	10	8	Q92XV3	Q9zyv3 diposaurus
43	13	23.6	10	8	Q92IV0	Q9zyv0 petrosaurus
44	13	23.6	10	8	Q92YU7	Q9zyu7 sator angus
45	13	23.6	10	8	Q92YU4	Q9zyu4 sceloporu
46	13	23.6	10	8	Q92YU1	Q9zyu1 uma scopari
47	13	23.6	10	8	Q92VT8	Q9zyt8 urosaurus g
48	13	23.6	10	8	Q92VT5	Q9zyt5 uta stansbu
49	13	23.6	10	8	Q92XS9	Q9zyS9 phymaturus
50	13	23.6	10	8	Q9TG98	Q9tG98 shinisaurus

ALIGNMENTS

RESULT 1

ID	P82440	PRELIMINARY;	PRT;	9 AA.
AC	P82440;			
DT	01-JUN-2000 (TREMBlrel. 14, Created)			
DT	01-JUN-2000 (TREMBlrel. 14, Last sequence update)			
DT	01-JUN-2000 (TREMBlrel. 14, Last annotation update)			
DE	42 KDA CELL WALL PROTEIN (FRAGMENT).			
OS	Nicotiana tabacum (Common tobacco).			
OC	Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;			
OC	Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;			
OC	Asteridae; euasterids I; Solanales; Solanaceae; Nicotiana.			
OX	NCBI_TaxID=4097;			
RP	[1]			
RP	SEQUENCE.			
RC	STRAIN=CV. PETIT HAVANA;			
RA	Blee K.A., Bonham V.A., Mitchell G.P., Robertson D., Slabas A.R.,			
RA	Wojtaszek P., Bolwell G.P.;			
RT	"Proteomic study of secondary cell wall proteins from transformed tobacco culture.";			
RL	Planta 0:0-0(2000).			
CC	-!- SUBCELLULAR LOCATION: CELL WALL.			
CC	-!- TISSUE SPECIFICITY: XYLEM.			
KW	Cell wall.			
FT	NON_TER			
SQ	SEQUENCE 9 AA; 1053 MW; 298CC9D2D5BB1B07 CRC64;			

Query Match 38.2%; Score 21; DB 10; Length 9;

Best Local Similarity 57.1%; Pred.No. 5.6e+05;

Matches 4; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 3 QKLFFFA 9

Db 3 EESVFFA 9

```
RESULT 2
Q9GD36 PRELIMINARY; PRT; 9 AA.
AC Q9GD36;
DT 01-MAR-2001 (TRENBLrel. 16, Created)
DT 01-MAR-2001 (TRENBLrel. 16, Last sequence update)
DT 01-MAR-2001 (TRENBLrel. 16, Last annotation update)
DE RIBOSOMAL PROTEIN S16 (FRAGMENT).
GN RPS16.
OS Juncus effusus.
OG Chloroplast.
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; Liliopsida; Poales; Juncaceae; Juncus.
OX NCBI_TaxID=13579;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=LEAF;
RA Asmussen C.B., Chase M.W.;
RT "Coding and noncoding plastid DNA in palm systematics.";
RL Submitted (JUL-2000) to the EMBL/GenBank/DDBJ databases.
DR EMBL; AJ404962; CAC17904.1; -.
KW Chloroplast.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1135 MW; 8DCCC9D2C046CB41 CRC64;

Query Match 34.5%; Score 19; DB 8; Length 9;
Best Local Similarity 60.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 4 KLVFF 8
DB 4 QIVFF 8

RESULT 3
Q924N8 PRELIMINARY; PRT; 9 AA.
AC Q924N8;
DT 01-DEC-2001 (TRENBLrel. 19, Created)
DT 01-DEC-2001 (TRENBLrel. 19, Last sequence update)
DT 01-DEC-2001 (TRENBLrel. 19, Last annotation update)
DE NIEMANN PICK TYPE C1 PROTEIN (FRAGMENT).
GN NPC1.
OS Mus musculus (Mouse).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Sclurognathi; Muridae; Murinae; Mus.
OX NCBI_TaxID=10090;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=C57BLKS;
RA Gevry N.Y., Lacroix D.A., Murphy B.D.;
RT "Niemann-Pick C1 protein gene, partial cds and promotor region.";
RL Submitted (SEP-1999) to the EMBL/GenBank/DDBJ databases.
DR EMBL; AF184964; AAK83683.1; -.
FT NON_TER 9 9
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 890 MW; 2C4E2DC761E1EDD8 CRC64;

Query Match 32.7%; Score 18; DB 11; Length 9;
Best Local Similarity 60.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHOKL 5
DB 4 HHPAL 8

RESULT 4
Q39952 PRELIMINARY; PRT; 10 AA.
ID Q39952
AC Q39952;

DT 01-JAN-1998 (TRENBLrel. 05, Created)
DT 01-JAN-1998 (TRENBLrel. 05, Last sequence update)
DT 01-DEC-2001 (TRENBLrel. 19, Last annotation update)
DE E1 PROTEIN (FRAGMENT).
OS Hepatitis GB virus C.
OC Viruses; ssRNA positive-strand viruses, no DNA stage; Flaviviridae;
OC GBV-C/HGV group.
OX NCBI_TaxID=39839;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=ED_INBURGH HAEMOPHILIAC;
RX MEDLINE=97368412; PubMed=9225026;
RA Smith D.B., Cuccenu N., Davidson F., Jarvis L.M., Mokili J.L.,
RA Hamid S., Ludlam C.A., Simmonds P.;
RT "Discrimination of hepatitis G virus/GBV-C geographical variants by
RT analysis of the 5' non-coding region.";
RL J. Gen. Virol. 78:1533-1542(1997).
DR EMBL; AF003170; AAC57981.1; -.
FT NON_TER 10 10
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1152 MW; CC88F0C9C7272732 CRC64;

Query Match 30.9%; Score 17; DB 12; Length 10;
Best Local Similarity 75.0%; Pred. No. 5.8e+03;
Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8
DB 5 LLFF 8

RESULT 5
Q9WLE4 PRELIMINARY; PRT; 10 AA.
ID Q9WLE4;
AC Q9WLE4;
DT 01-NOV-1999 (TRENBLrel. 12, Created)
DT 01-NOV-1999 (TRENBLrel. 12, Last sequence update)
DT 01-DEC-2001 (TRENBLrel. 19, Last annotation update)
DE POLYPROTEIN (FRAGMENT).
OS Hepatitis G virus.
OC Viruses; ssRNA positive-strand viruses, no DNA stage; Flaviviridae;
OC GBV-C/HGV group.
OX NCBI_TaxID=45255;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=SG3403;
RX MEDLINE=99266893; PubMed=10335862;
RA Wong S.B.J., Chan S.H., Ren E.C.;
RT "Diversity of GB virus C/hepatitis G virus isolates in Singapore:
RT predominance of group 2a and the Asian group 3 variant.";
RL J. Med. Virol. 58:145-153(1999).
DR EMBL; AF078063; AAC32369.1; -.
FT NON_TER 10 10
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1152 MW; CC88F0C9C7272732 CRC64;

Query Match 30.9%; Score 17; DB 12; Length 10;
Best Local Similarity 75.0%; Pred. No. 5.8e+03;
Matches 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8
DB 5 LLFF 8

RESULT 6
P82070 PRELIMINARY; PRT; 5 AA.
ID P82070
AC P82070;
DT 01-MAY-2000 (TRENBLrel. 13, Created)
DT 01-MAY-2000 (TRENBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TRENBLrel. 13, Last annotation update)
DE RUBELLIDIN 1.1.
OS Litoria rubella (Desert tree frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
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OC Amphibia; Batrachia; Anura; Neobatrachia; Bufonoidea; Hylidae;
OC Litoria.
OX NCBI_TaxID=104895;
RN [1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=SKIN SECRETION;
RA Steinborner S.T., Wabnitz P.A., Waugh R.J., Bowie J.H., Gao C.,
RA Tyler M.J., Wallace J.C.;
RT 'The structure of new peptides from the Australian red tree frog
RT 'Litoria rubella'. the skin peptide profile as a probe for the study
RT of evolutionary trends of amphibians.';
RL Aust. J. Chem. 49:955-963(1996).
CC -1- FUNCTION: CAERIDINS SHOW NEITHER NEUROPEPTIDE ACTIVITY NOR
CC ANTIBIOTIC ACTIVITY.
CC -1- TISSUE SPECIFICITY: SECRETED BY THE SKIN DORSAL GLANDS.
CC -1- MASS SPECTROMETRY: MW=598; METHOD=FAB.
KW Amphibian skin.
SQ SEQUENCE 5 AA; 598 MW; 6D9C9CAB2A00000 CRC64;

Query Match 29.1%; Score 16; DB 13; Length 5;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
Db 3 FFA 5

RESULT 7
P82071 ID P82071 PRELIMINARY; PRT; 5 AA.
AC P82071;
DT 01-MAY-2000 (TREMBlrel. 13, Created)
DT 01-MAY-2000 (TREMBlrel. 13, Last sequence update)
DT 01-MAY-2000 (TREMBlrel. 13, Last annotation update)
DE RUBELLIDIN 2.1.
OS Litoria rubella (Desert tree frog).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Amphibia; Batrachia; Anura; Neobatrachia; Bufonoidea; Hylidae;
OC Litoria.
OX NCBI_TaxID=104895;
RN [1]
RP SEQUENCE, AND MASS SPECTROMETRY.
RC TISSUE=SKIN SECRETION;
RA Steinborner S.T., Wabnitz P.A., Waugh R.J., Bowie J.H., Gao C.,
RA Tyler M.J., Wallace J.C.;
RT 'The structure of new peptides from the Australian red tree frog
RT 'Litoria rubella'. the skin peptide profile as a probe for the study
RT of evolutionary trends of amphibians.';
RL Aust. J. Chem. 49:955-963(1996).
CC -1- FUNCTION: CAERIDINS SHOW NEITHER NEUROPEPTIDE ACTIVITY NOR
CC ANTIBIOTIC ACTIVITY.
CC -1- TISSUE SPECIFICITY: SECRETED BY THE SKIN DORSAL GLANDS.
CC -1- MASS SPECTROMETRY: MW=626; METHOD=FAB.
KW Amphibian skin.
SQ SEQUENCE 5 AA; 626 MW; 6D9C9CB10300000 CRC64;

Query Match 29.1%; Score 16; DB 13; Length 5;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 7 FFA 9
Db 3 FFA 5

RESULT 8
Q15894 ID Q15894 PRELIMINARY; PRT; 8 AA.
AC Q15894;
DT 01-NOV-1996 (TREMBlrel. 01, Created)
DT 01-NOV-1996 (TREMBlrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)

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DE (CLONE XP587B) (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=PLACENTA;
RA Lee C.-C., Yazdani A., Wehnert M., Bailey J., Couch L., Xiong M.,
RA Coolbaugh M.I., Chnault C.A., Baidini A., Lindsay E.A., Zhao Z.-Y.,
RA Caskey C.T.H.;
RT "Isolation of chromosome-specific genes by reciprocal probing of
RT arrayed cDNAs and cosmid libraries.";
RL Hum. Mol. Genet. 0:0-0(1995).
DR EMBL: L32074; AAA73884.1; -.
ET NON_TER 1 1
FT NON_TER 8 8
SQ SEQUENCE 8 AA; 952 MW; EBC735B1E1F1B6D6 CRC64;

Query Match 29.1%; Score 16; DB 4; Length 8;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HH 2
Db 4 HH 5

RESULT 9
Q40530 ID Q40530 PRELIMINARY; PRT; 8 AA.
AC Q40530;
DT 01-NOV-1996 (TREMBlrel. 01, Created)
DT 01-NOV-1996 (TREMBlrel. 01, Last sequence update)
DT 01-NOV-1998 (TREMBlrel. 08, Last annotation update)
DE P20 N WITH A LEADER PEPTIDE.
OS Nicotiana tabacum (Common tobacco).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots;
OC Asteridae; euasterids I; Solanales; Solanaceae; Nicotiana.
OX NCBI_TaxID=4097;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=87089808; PubMed=3540612;
RA Herman L.M.F., Montagu M.C.V., Depicker A.G.;
RT "Isolation of tobacco DNA segments with plant promoter activity.";
RL Mol. Cell. Biol. 6:4486-4492(1986).
DR EMBL: M14685; AAA34090.1; -.
SQ SEQUENCE 8 AA; 1109 MW; E257205B19C9C6 CRC64;

Query Match 29.1%; Score 16; DB 10; Length 8;
Best Local Similarity 60.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

Qy 6 VFFAE 10
Db 1 MEFFE 5

RESULT 10
Q47556 ID Q47556 PRELIMINARY; PRT; 9 AA.
AC Q47556;
DT 01-NOV-1996 (TREMBlrel. 01, Created)
DT 01-NOV-1996 (TREMBlrel. 01, Last sequence update)
DT 01-NOV-1998 (TREMBlrel. 08, Last annotation update)
DE ASPARTATE TRANSCARBAMOYLASE REGULATORY CHAIN (FRAGMENT).
GN PYRI.
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OX NCBI_TaxID=562;
RN [1]

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RP SEQUENCE FROM N.A.
RX MEDLINE=82275057; PubMed=7051000;
RA Pauza C.D., Karels M.J., Navre M., Schachman H.K.;
RT "Genes encoding Escherichia coli aspartate transcarbamoylase: The
PYR-pyri operon.";
RL Proc. Natl. Acad. Sci. U.S.A. 79:4020-4024(1982).
RN (2)
RP SEQUENCE OF 1-5 FROM N.A.
RX MEDLINE=83195078; PubMed=6302686;
RA Hoover T.A., Roof W.D., Foltermann K.F., O'Donovan G.A., Bencini D.A.,
RA Wild J.R.;
RT "Nucleotide sequence of the structural gene (pyrB) that encodes the
RT catalytic polypeptide of aspartate transcarbamoylase of Escherichia
RT coli.";
RL Proc. Natl. Acad. Sci. U.S.A. 80:2462-2466(1983).
DR EMBL: J01670; AAA24475.1; -.
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1085 MW; 99EFD723344AA1F1 CRC64;

Query Match 29.1%; Score 16; DB 2; Length 9;
Best Local Similarity 60.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHOKL 5
| | |
Db 3 HDNKL 7

RESULT 11
Q9H4M8 PRELIMINARY; PRT; 9 AA.
ID Q9H4M8
AC Q9H4M8
DT 01-MAR-2001 (TrEMBLrel. 16, Created)
DT 01-MAR-2001 (TrEMBLrel. 16, Last sequence update)
DT 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
DE PAR2 (FRAGMENT).
GN NR112.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=PERIPHERAL BLOOD;
RA Pentecost B.T., Ling G.;
RT "The human pregnane X receptor promoter complex provides
RT transcriptional starts for a number of PXR related transcripts.";
RL Submitted (AUG-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL: AY007189; AAG23345.1; -.
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1129 MW; 82F8E1F1B411B2D1 CRC64;

Query Match 29.1%; Score 16; DB 4; Length 9;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HH 2
| |
Db 7 HH 8

RESULT 12
Q9R5L7 PRELIMINARY; PRT; 8 AA.
ID Q9R5L7
AC Q9R5L7
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT 01-JUN-2000 (TrEMBLrel. 14, Last annotation update)
DE 1,4-BETA-D-GLUCAN GLUCANOHYDROLASE (EC 3.2.1.4) (FRAGMENT).
OS Clostridium thermocellum.
OC Bacteria; Firmicutes; Bacillus/Clostridium group; Clostridiaceae;
OC Clostridium
OX NCBI_TaxID=1515;

RN SEQUENCE.
RX MEDLINE=92231850; PubMed=1567379;
RA Romaniec M.P., Fauth U., Kobayashi T., Huskisson N.S., Barker P.J.,
RA Demain A.L.;
RT "Purification and characterization of a new endoglucanase from
RT Clostridium thermocellum.";
RL Biochem. J. 283:69-73(1992).
SQ SEQUENCE 8 AA; 823 MW; C2CIAB1DD9D1B775 CRC64;

Query Match 27.3%; Score 15; DB 2; Length 8;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
| | |
Db 4 FAE 6

RESULT 13
O13591 PRELIMINARY; PRT; 8 AA.
ID O13591
AC O13591
DT 01-JAN-1998 (TrEMBLrel. 05, Created)
DT 01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
DT 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
DE ORF YNL337W (FRAGMENT).
GN YNL337W.
OS Saccharomyces cerevisiae (Baker's yeast).
OC Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
OC Saccharomycetales; Saccharomycetaceae; Saccharomycetes.
OX NCBI_TaxID=4932;
RN [1]
RP SEQUENCE FROM N.A.
RA Obermaier B., Piravandi E., Rinke M.;
RL Submitted (MAY-1996) to the EMBL/GenBank/DBJ databases.
RN [2]
RP SEQUENCE FROM N.A.
RA MIPS;
RL Submitted (APR-1996) to the EMBL/GenBank/DBJ databases.
DR EMBL: 271612; CAA96271.2; -.
DR SGD: S0005281; YNL337W.
FT NON_TER 1 1
SQ SEQUENCE 8 AA; 1005 MW; 5CA441E449C9C720 CRC64;

Query Match 27.3%; Score 15; DB 3; Length 8;
Best Local Similarity 50.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8
: | | |
Db 1 ILFF 4

RESULT 14
Q15889 PRELIMINARY; PRT; 8 AA.
ID Q15889
AC Q15889
DT 01-NOV-1996 (TrEMBLrel. 01, Created)
DT 01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE (CLONE XP15H8B) (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=PLACENTA;
RA Lee C.-C., Yazdani A., Wehnert M., Bailey J., Couch L., Xiong M.,
RA Coolbaugh M.I., Chinault C.A., Baldini A., Lindsay E.A., Zhao Z.-Y.,
RA Caskey C.T.H.;
RT "Isolation of chromosome-specific genes by reciprocal probing of


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Query Match      27.3%; Score 15; DB 12; Length 9;
Best Local Similarity 50.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKL 5
DB 2 HNKI 5

RESULT 19
Q9R7J8 PRELIMINARY; PRT; 10 AA.
ID Q9R7J8
AC Q9R7J8;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DE 01-JUN-2000 (TrEMBLrel. 14, Last annotation update)
DE VACUOLATING CYTOTOXIN (FRAGMENT).
GN VACA.
OS Helicobacter pylori (Campylobacter pylori).
OC Bacteria; Proteobacteria; epsilon subdivision; Helicobacter group;
OC Helicobacter.
OX NCBI_TaxID=210;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN-KOBE 500;
RA Shirasaka D.;
RT "Helicobacter pylori vacA gene, strain Kobe 500, partial cds.";
DR EMBL: AB017599; BAA33412.1; -.
FT NON_TER 1
FT NON_TER 10
SQ SEQUENCE 10 AA; 1018 MW; 414390C76879CDD7 CRC64;

Query Match      27.3%; Score 15; DB 2; Length 10;
Best Local Similarity 75.0%; Pred. No. 1.4e+04;
Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 4 KLVE 7
DB 2 KLAF 5

RESULT 20
Q9UCS3 PRELIMINARY; PRT; 10 AA.
ID Q9UCS3
AC Q9UCS3;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TrEMBLrel. 13, Last annotation update)
DE TROPOMYOSIN-33 KDA CALCIUM BINDING PROTEIN FRAGMENT D.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE.
RX MEDLINE=92090441; PubMed=1836432;
RA Crabos M., Yamakado T., Heizmann C.W., Cerletti N., Buhler F.R.,
RA Erne P.;
RT "the calcium binding protein tropomyosin in human platelets and
RT cardiac tissue: elevation in hypertensive cardiac hypertrophy.";
RL Eur. J. Clin. Investig. 21:472-478(1991).
SQ SEQUENCE 10 AA; 1126 MW; 7A44FD3DC2DAFAEB CRC64;

Query Match      27.3%; Score 15; DB 4; Length 10;
Best Local Similarity 100.0%; Pred. No. 1.4e+04;
Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 8 FAE 10
DB 1 FAE 3

Query Match      27.3%; Score 15; DB 5; Length 10;
Best Local Similarity 40.0%; Pred. No. 1.4e+04;
Matches 2; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKLV 6
DB 6 HSKVL 10

RESULT 21
P82223 PRELIMINARY; PRT; 10 AA.
ID P82223
AC P82223;
DT 01-OCT-2001 (TrEMBLrel. 18, Created)
DT 01-OCT-2001 (TrEMBLrel. 18, Last sequence update)
DE UNKNOWN PROTEIN FROM 2D-PAGE (FRAGMENT).
OS Bombyx mori (Silk moth).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia;
OC Bombycoidea; Bombycidae; Bombyx.
OX NCBI_TaxID=7091;
RN [1]
RP SEQUENCE.
RC STRAIN-XINHANG X KEMING; TISSUE=BODY WALL, AND FAT BODY;
RX MEDLINE=21177481; PubMed=11280994;
RA Zhong B.X.;
RT "Protein database for several tissues derived from five instar of
RT silkworm.";
RL I Chuan Hsueh Pao 28:217-224(2001).
FT NON_TER 10
FT NON_TER 10
SQ SEQUENCE 10 AA; 1054 MW; D0F722C325B1F1B2 CRC64;

Query Match      27.3%; Score 15; DB 5; Length 10;
Best Local Similarity 40.0%; Pred. No. 1.4e+04;
Matches 2; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKLV 6
DB 5 HSKVL 9

RESULT 22
P82224 PRELIMINARY; PRT; 10 AA.
ID P82224
AC P82224;
DT 01-OCT-2001 (TrEMBLrel. 18, Created)
DT 01-OCT-2001 (TrEMBLrel. 18, Last sequence update)
DT 01-OCT-2001 (TrEMBLrel. 18, Last annotation update)
DE UNKNOWN PROTEIN FROM 2D-PAGE (FRAGMENT).
OS Bombyx mori (Silk moth).
OC Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
OC Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia;
OC Bombycoidea; Bombycidae; Bombyx.
OX NCBI_TaxID=7091;
RN [1]
RP SEQUENCE.
RC STRAIN-XINHANG X KEMING; TISSUE=BODY WALL, AND FAT BODY;
RX MEDLINE=21177481; PubMed=11280994;
RA Zhong B.X.;
RT "Protein database for several tissues derived from five instar of
RT silkworm.";
RL I Chuan Hsueh Pao 28:217-224(2001).
FT NON_TER 10
FT NON_TER 10
SQ SEQUENCE 10 AA; 1064 MW; D77CBE25B1F1B2CD CRC64;

Query Match      27.3%; Score 15; DB 5; Length 10;
Best Local Similarity 40.0%; Pred. No. 1.4e+04;
Matches 2; Conservative 2; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKLV 6
DB 6 HSKVL 10

RESULT 23
Q9UMH9 PRELIMINARY; PRT; 8 AA.
ID Q9UMH9
AC Q9UMH9;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TrEMBLrel. 13, Last annotation update)

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DE RHCE PROTEIN (FRAGMENT).
GN RHCE.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=BLOOD;
RX MEDLINE=97260406; PubMed=9106526;
RA Matassi G., Cherif-Zahar B., Moura I., Cartron J.P.;
RT "Characterization of the recombination hot spot involved in the
RT genomic rearrangement leading to the hybrid D-CE-D gene in the DVI
RT phenotype.";
RL Am. J. Hum. Genet. 60:808-817(1997).
RN [2]
RP SEQUENCE FROM N.A.
RC TISSUE=BLOOD;
RX MEDLINE=90349591; PubMed=1696722;
RA Cherif-Zahar B., Bloy C., Le Van Kim C., Blanchard D., Bailly P.,
RA Hermand P., Salmon C., Cartron J.-P., Colin Y.;
RT "Molecular cloning and protein structure of a human blood group Rh
RT polypeptide.";
RL Proc. Natl. Acad. Sci. U.S.A. 87:6243-6247(1990).
DR EMBL; Z97030; CAB09726.1; -.
FT NON_TER 1
FT NON_TER 8
SQ SEQUENCE 8 AA; 1049 MW; C007244691FB5AB1 CRC64;

Query Match 25.5%; Score 14; DB 4; Length 8;
Best Local Similarity 40.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHQKL 5
DB 3 YHML 7

RESULT 24
Q9UC36 PRELIMINARY; PRT; 9 AA.
AC Q9UC36;
DT 01-MAY-2000 (TReMBLrel. 13, Created)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TReMBLrel. 13, Last annotation update)
DE 28 KDA HEAT SHOCK PROTEIN HOMOLOG FRAGMENT 1.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE.
RX MEDLINE=92218434; PubMed=1560006;
RA Kato K., Shinohara H., Goto S., Inaguma Y., Morishita R., Asano T.;
RT "Copurification of small heat shock protein with alpha B crystallin
RT from human skeletal muscle.";
RL J. Biol. Chem. 267:7718-7725(1992).
SQ SEQUENCE 9 AA; 1220 MW; 26933415B1F77B43 CRC64;

Query Match 25.5%; Score 14; DB 4; Length 9;
Best Local Similarity 50.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 2 HQKL 5
DB 5 HSRL 8

RESULT 25
Q94VG2 PRELIMINARY; PRT; 9 AA.
ID Q94VG2
AC Q94VG2;
DT 01-DEC-2001 (TReMBLrel. 19, Created)

DT 01-DEC-2001 (TReMBLrel. 19, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Varanus indicus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Scleroglossa; Anguilliformia; Varanidae; Varanus.
OX NCBI_TaxID=62043;
RN [1]
RP SEQUENCE FROM N.A.
RA Ast J.C.;
RT "Mitochondrial DNA evidence and evolution in Varanoidea (Squamata).";
RL Cladistics 17:0-0(2001).
DR EMBL; AF407505; AAL10069.1; -.
KW Mitochondrion.
FT NON_TER 9
SQ SEQUENCE 9 AA; 1258 MW; 881259C727336411 CRC64;

Query Match 25.5%; Score 14; DB 8; Length 9;
Best Local Similarity 50.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 2; Mismatches 0; Indels 0; Gaps 0;

QY 5 LVFF 8
DB 5 LLFY 8

RESULT 26
P82937 PRELIMINARY; PRT; 10 AA.
AC P82937
DT 01-MAR-2001 (TReMBLrel. 16, Created)
DT 01-MAR-2001 (TReMBLrel. 16, Last sequence update)
DT 01-JUN-2001 (TReMBLrel. 17, Last annotation update)
DE UNKNOWN ENDOSPERM PROTEIN B (FRAGMENT).
OS Hordeum vulgare (Barley).
OC Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
OC Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Poaceae;
OC Triticeae; Hordeum.
OX NCBI_TaxID=4513;
RN [1]
RP SEQUENCE.
RC STRAIN=CV. BOWI; TISSUE=STARCHY ENDOSPERM;
RX MEDLINE=21088911; PubMed=11271488;
RA Kristoffersen H.E., Flengsrud R.;
RT "Separation and characterization of basic barley seed proteins.";
RL Electrophoresis 21:3693-3700(2000).
CC -!- MISCELLANEOUS: ON THE 2D-GEL THE DETERMINED PI OF THIS UNKNOWN
CC PROTEIN IS: 8.5-9.0, ITS MW IS: 11.9 KDA.
FT NON_TER 10
FT NON_TER 10
SQ SEQUENCE 10 AA; 1297 MW; 8248A50B11FB5EBA CRC64;

Query Match 25.5%; Score 14; DB 10; Length 10;
Best Local Similarity 25.0%; Pred. No. 2.2e+04;
Matches 1; Conservative 3; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQK 4
DB 5 YHER 8

RESULT 27
O09258 PRELIMINARY; PRT; 8 AA.
ID O09258
AC O09258;
DT 01-JUL-1997 (TReMBLrel. 04, Created)
DT 01-JUL-1997 (TReMBLrel. 04, Last sequence update)
DT 01-DEC-2001 (TReMBLrel. 19, Last annotation update)
DE NIFH (FRAGMENT).
GN NIFH.
OS Synechococcus sp. (strain PCC 8801 / RF-1) (Cyanothecae PCC 8801).
OC Bacteria; Cyanobacteria; Chroococcales; Cyanothecae.

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OX NCBI_TaxID=41431;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=RF-1;
RX MEDLINE=99231861; PubMed=10217509;
RA Huang T.C., Lin R.F., Chu M.K., Chen H.M.;
RT "Organization and expression of nitrogen-fixation genes in the aerobic
RT nitrogen-fixing unicellular cyanobacterium Synechococcus sp. strain
RT RF-1."
RL Microbiology 145:743-753(1999).
DR EMBL: AF001780; AAC33369.1; -.
FT NON_TER 8
SQ SEQUENCE 8 AA; 985 MW; F16B59CDD046C406 CRC64;

Query Match 23.6%; Score 13; DB 2; Length 8;
Best Local Similarity 16.7%; Pred. No. 5.6e+05;
Matches 1; Conservative 4; Mismatches 1; Indels 0; Gaps 0;

QY 3 QKLVFF 8
   : : :
Db 2 RQIAFY 7

RESULT 28
Q9S6D5 PRELIMINARY; PRT; 8 AA.
AC Q9S6D5;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TrEMBLrel. 13, Last annotation update)
DE PUTATIVE IS30 TRANSPOSASE (FRAGMENT).
OS Escherichia coli.
OC Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Escherichia.
OC NCBI_TaxID=562;
OX [1]
RN SEQUENCE FROM N.A.
RP STRAIN=A295B;
RX MEDLINE=99194747; PubMed=10094716;
RA Rahn A., Drummelsmith J., Whitfield C.;
RT "Conserved organization in the cps gene clusters for expression of
RT Escherichia coli group 1 k antigens: relationship to the colanic acid
RT biosynthesis locus and the cps genes from Klebsiella pneumoniae.";
RL J. Bacteriol. 181:2307-2313(1999).
DR EMBL: AF118251; AAD30008.1; -.
FT NON_TER 8
SQ SEQUENCE 8 AA; 1011 MW; F21DC1A9D1B41406 CRC64;

Query Match 23.6%; Score 13; DB 2; Length 8;
Best Local Similarity 16.7%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 7 FFAE 10
   : : :
Db 5 FTAE 8

RESULT 29
Q56759 PRELIMINARY; PRT; 8 AA.
AC Q56759;
DT 01-NOV-1996 (TrEMBLrel. 01, Created)
DT 01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE HALOACID DEHALOGENASE (FRAGMENT).
GN DHLB.
OS Xanthobacter autotrophicus.
OC Bacteria; Proteobacteria; alpha subdivision; Rhizobiaceae group;
OC Hyphomicrobium group; Xanthobacter.
OX NCBI_TaxID=280;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=GJ10, AND CV. M50;

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RX MEDLINE=95173113; PubMed=7868610;
RA Van der Ploeg J., Willemssen M., van Hall G., Janssen D.B.;
RT "Adaptation of xanthobacter autotrophicus GJ10 to bromoacetate due to
RT activation and mobilization of the haloacetate dehalogenase gene by
RT insertion element IS1247."
RL J. Bacteriol. 177:1348-1356(1995).
DR EMBL: X84038; CAA58857.1; -.
FT NON_TER 8
SQ SEQUENCE 8 AA; 922 MW; F3A9D2D2CDD33056 CRC64;

Query Match 23.6%; Score 13; DB 2; Length 8;
Best Local Similarity 60.0%; Pred. No. 5.6e+05;
Matches 3; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 4 KLVFF 8
   : : :
Db 3 KAVVF 7

RESULT 30
Q9UDZ4 PRELIMINARY; PRT; 8 AA.
AC Q9UDZ4;
DT 01-MAY-2000 (TrEMBLrel. 13, Created)
DT 01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE RHD PROTEIN (FRAGMENT).
GN RHD.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=BLOOD;
RX MEDLINE=97260406; PubMed=9106526;
RA Matassi G., Cherif-Zahar B., Mouro I., Cartron J.P.;
RT "Characterization of the recombination hot spot involved in the
RT genomic rearrangement leading to the hybrid D-CE-D gene in the DVI
RT phenotype."
RL Am. J. Hum. Genet. 60:808-817(1997).
RN [2]
RP SEQUENCE FROM N.A.
RC TISSUE=BLOOD;
RX MEDLINE=93066356; PubMed=1438298;
RA Le Van Kim C., Mouro I., Cherif-Zahar B., Raynal V., Cherrier C.,
RA Cartron J.P., Colin Y.;
RT "Molecular cloning and primary structure of the human blood group Rh
RT polypeptide."
RL Proc. Natl. Acad. Sci. U.S.A. 89:10925-10929(1992).
DR EMBL: Z97031; CAB09727.1; -.
FT NON_TER 1
FT NON_TER 8
SQ SEQUENCE 8 AA; 1042 MW; D296944691FB5AB1 CRC64;

Query Match 23.6%; Score 13; DB 4; Length 8;
Best Local Similarity 16.7%; Pred. No. 5.6e+05;
Matches 1; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

QY 1 HHOKLV 6
   : : :
Db 3 YHNMNM 8

RESULT 31
O15899 PRELIMINARY; PRT; 8 AA.
AC O15899;
DT 01-JAN-1998 (TrEMBLrel. 05, Created)
DT 01-JAN-1998 (TrEMBLrel. 05, Last sequence update)
DT 01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
DE 12D3 ANTIGEN (FRAGMENT).
GN B012D3.

```


OS Babesia ovis.
 OC Eukaryota; Alveolata; Apicomplexa; Piroplasmida; Babesiidae; Babesia.
 OX NCBI_TaxID=5869;
 RN [1]
 RP SEQUENCE FROM N.A.
 RC STRAIN=ANKARA;
 RA Silins G.U., Blakeley R.L., Riddles P.W.;
 RT "Characterization of the transpositional control region of the 12D3
 RT antigen gene from the sporozoan Babesia bovis.";
 RL Submitted (JAN-1996) to the EMBL/GenBank/DBJ databases.
 DR EMBL; U49199; AAB66365.1; -.
 FT NON_TER 8
 SQ SEQUENCE 8 AA; 992 MW; F0C7273411B2C726 CRC64;
 Query Match 23.6%; Score 13; DB 5; Length 8;
 Best Local Similarity 50.0%; Pred. No. 5.6e+05;
 Matches 2; Conservative 2; Mismatches 0; Indels 0; Gaps 0;
 QY 4 KLVF 7
 Db :I:I
 5 RLLF 8
 RESULT 32
 Q9GMH3 PRELIMINARY; PRT; 8 AA.
 AC Q9GMH3
 DT 01-MAR-2001 (TREMBLrel. 16, Created)
 DT 01-MAR-2001 (TREMBLrel. 16, Last sequence update)
 DE ACTIN (FRAGMENT).
 OS Lagenorhynchus obscurus (dusky dolphin).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
 OC Lagenorhynchus.
 OX NCBI_TaxID=27611;
 RN [1]
 RP SEQUENCE FROM N.A.
 RA Hare M.P., Cipriano F., Palumbi S.R.;
 RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
 RT Speciation, Systematics and Conservation.";
 RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
 DR EMBL; AF140833; AAF98686.1; -.
 FT NON_TER 1
 FT NON_TER 8
 SQ SEQUENCE 8 AA; 962 MW; 5BD1F417740862C0 CRC64;
 Query Match 23.6%; Score 13; DB 6; Length 8;
 Best Local Similarity 100.0%; Pred. No. 5.6e+05;
 Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2 HQ 3
 Db :I:I
 7 HQ 8
 RESULT 33
 Q28866 PRELIMINARY; PRT; 8 AA.
 AC Q28866
 DT 01-NOV-1996 (TREMBLrel. 01, Created)
 DT 01-NOV-1996 (TREMBLrel. 01, Last sequence update)
 DT 01-MAY-1999 (TREMBLrel. 10, Last annotation update)
 DE ACTIN PROTEIN (FRAGMENT).
 OS Megaptera novaeangliae (Humpback whale).
 GN Megaptera novaeangliae (Humpback whale).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Mysticeti;
 OC Balaenopteridae; Megaptera.
 OX NCBI_TaxID=9773;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=94285813; PubMed=7912407;

RA Palumbi S.R., Baker C.S.;
 RT "Contrasting population structure from nuclear intron sequences and
 RT mtDNA of humpback whales.";
 RL Mol. Biol. Evol. 11:426-435(1994).
 DR EMBL; S73467; AAD14118.1; -.
 FT NON_TER 1
 SQ SEQUENCE 8 AA; 906 MW; 69C866D1F4177408 CRC64;
 Query Match 23.6%; Score 13; DB 6; Length 8;
 Best Local Similarity 100.0%; Pred. No. 5.6e+05;
 Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 2 HQ 3
 Db :I:I
 5 HQ 6
 RESULT 34
 O02831 PRELIMINARY; PRT; 8 AA.
 AC O02831
 DT 01-JUL-1997 (TREMBLrel. 04, Created)
 DT 01-JUL-1997 (TREMBLrel. 04, Last sequence update)
 DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
 DE PRO ALPHA 1 TYPE III COLLAGEN PROTEIN (FRAGMENT).
 OS Oryctolagus cuniculus (Rabbit).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Lagomorpha; Leporidae; Oryctolagus.
 OX NCBI_TaxID=9986;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=96377339; PubMed=8783186;
 RA Metsaranta M., Kujala U.M., Pelliniemi L., Osterman H., Aho H.,
 RA Vuorio E.;
 RT "Evidence for insufficient chondrocytic differentiation during repair
 RT of full-thickness defects of articular cartilage.";
 RL Matrix Biol. 15:39-47(1996).
 DR EMBL; S83371; AAD14433.1; -.
 KW Collagen.
 FT NON_TER 1
 SQ SEQUENCE 8 AA; 1028 MW; B859C7272EA77371 CRC64;
 Query Match 23.6%; Score 13; DB 6; Length 8;
 Best Local Similarity 42.9%; Pred. No. 5.6e+05;
 Matches 3; Conservative 1; Mismatches 3; Indels 0; Gaps 0;
 QY 1 HHQKLVF 7
 Db :I:I
 1 HWPCLLF 7
 RESULT 35
 Q99NX9 PRELIMINARY; PRT; 8 AA.
 AC Q99NX9
 DT 01-JUN-2001 (TREMBLrel. 17, Created)
 DT 01-JUN-2001 (TREMBLrel. 17, Last sequence update)
 DT 01-DEC-2001 (TREMBLrel. 19, Last annotation update)
 DE AMYLOID BETA PROTEIN (FRAGMENT).
 GN APP.
 OS Hydrocoerus hydrochaeris (Capybara) (Carpincho).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Hystricognathi; Hydrochaeridae;
 OC Hydrochaeris.
 OX NCBI_TaxID=10149;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=21082082; PubMed=11214319;
 RA Murphy W.J., Eizirik E., Johnson W.E., Zhang Y.P., Ryder O.A.,
 RA O'Brien S.J.;
 RT "Molecular phylogenetics and the origins of placental mammals.";
 RL Nature 409:614-618(2001).
 DR EMBL; AY011342; AAG47377.1; -.

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FT NON_TER 1 1
SQ SEQUENCE 8 AA; 1071 MW; 1356D686DB19C9C3 CRC64;

Query Match
Best Local Similarity 23.6%; Score 13; DB 11; Length 8;
Matches 2; Conservative 1; Mismatches 1; Indels 0; Gaps 0;

QY 7 FFAE 10
DB 2 FFEQ 5

RESULT 36
Q15891 ID Q15891 PRELIMINARY; PRT; 9 AA.
AC Q15891;
DT 01-NOV-1996 (TREMBlrel. 01, Created)
DT 01-NOV-1996 (TREMBlrel. 01, Last sequence update)
DT 01-DEC-2001 (TREMBlrel. 19, Last annotation update)
DE (CLONE XP2E8B) (FRAGMENT).
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=PLACENTA;
RA Lee C.-C., Yazdani A., Wehnert M., Bailey J., Couch L., Xiong M.,
RA Coolbaugh M.I., Chinault C.A., Baldini A., Lindsay E.A., Zhao Z.-Y.,
RA Caskey C.T.H.;
RT Isolation of chromosome-specific genes by reciprocal probing of
RT arrayed cDNAs and cosmid libraries.
RL Hum. Mol. Genet. 0:0-0(1995).
DR EMBL; L32131; AAA73881.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1030 MW; E56635A1A33686D1 CRC64;

Query Match
Best Local Similarity 23.6%; Score 13; DB 4; Length 9;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQ 3
DB 2 HQ 3

RESULT 37
Q9GJV3 ID Q9GJV3 PRELIMINARY; PRT; 9 AA.
AC Q9GJV3;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE ACTIN (FRAGMENT).
OS Lagenorhynchus obscurus (dusky dolphin).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
OC Lagenorhynchus.
OX NCBI_TaxID=27611;
RN [1]
RP SEQUENCE FROM N.A.
RA Hare M.P., Cipriano F., Palumbi S.R.;
RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
RT Speciation, Systematics and Conservation.";
RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF140834; AAF98687.1; -.
DR EMBL; AF140832; AAF98685.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1049 MW; 1D0EF417740862C0 CRC64;

Query Match
Best Local Similarity 23.6%; Score 13; DB 6; Length 9;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQ 3
DB 2 HQ 3

RESULT 38
Q9GJV2 ID Q9GJV2 PRELIMINARY; PRT; 9 AA.
AC Q9GJV2;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE ACTIN (FRAGMENT).
OS Lagenorhynchus obliquidens (Pacific white-sided dolphin).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
OC Lagenorhynchus.
OX NCBI_TaxID=90247;
RN [1]
RP SEQUENCE FROM N.A.
RA Hare M.P., Cipriano F., Palumbi S.R.;
RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
RT Speciation, Systematics and Conservation.";
RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF140831; AAF98684.1; -.
DR EMBL; AF140826; AAF98679.1; -.
DR EMBL; AF140827; AAF98680.1; -.
DR EMBL; AF140828; AAF98681.1; -.
DR EMBL; AF140829; AAF98682.1; -.
DR EMBL; AF140830; AAF98683.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1049 MW; 1D0EF417740862C0 CRC64;

Query Match
Best Local Similarity 23.6%; Score 13; DB 6; Length 9;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQ 3
DB 8 HQ 9

RESULT 39
Q9GJV1 ID Q9GJV1 PRELIMINARY; PRT; 9 AA.
AC Q9GJV1;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE ACTIN (FRAGMENT).
OS Lagenorhynchus acutus (Atlantic white-sided dolphin).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
OC Lagenorhynchus.
OX NCBI_TaxID=90246;
RN [1]
RP SEQUENCE FROM N.A.
RA Hare M.P., Cipriano F., Palumbi S.R.;
RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
RT Speciation, Systematics and Conservation.";
RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF140825; AAF98678.1; -.
DR EMBL; AF140822; AAF98675.1; -.
DR EMBL; AF140823; AAF98676.1; -.
DR EMBL; AF140824; AAF98677.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1049 MW; 1D0EF417740862C0 CRC64;

Query Match
Best Local Similarity 23.6%; Score 13; DB 6; Length 9;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQ 3
DB 8 HQ 9

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Best Local Similarity 100.0%; Pred. No. 5.6e+05; Indels 0; Gaps 0;
Matches 2; Conservative 0; Mismatches 0;

QY 2 HQ 3
DB 8 HQ 9

RESULT 38
Q9GJV2 ID Q9GJV2 PRELIMINARY; PRT; 9 AA.
AC Q9GJV2;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE ACTIN (FRAGMENT).
OS Lagenorhynchus obliquidens (Pacific white-sided dolphin).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
OC Lagenorhynchus.
OX NCBI_TaxID=90247;
RN [1]
RP SEQUENCE FROM N.A.
RA Hare M.P., Cipriano F., Palumbi S.R.;
RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
RT Speciation, Systematics and Conservation.";
RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF140831; AAF98684.1; -.
DR EMBL; AF140826; AAF98679.1; -.
DR EMBL; AF140827; AAF98680.1; -.
DR EMBL; AF140828; AAF98681.1; -.
DR EMBL; AF140829; AAF98682.1; -.
DR EMBL; AF140830; AAF98683.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1049 MW; 1D0EF417740862C0 CRC64;

Query Match
Best Local Similarity 100.0%; Pred. No. 5.6e+05; Indels 0; Gaps 0;
Matches 2; Conservative 0; Mismatches 0;

QY 2 HQ 3
DB 8 HQ 9

RESULT 39
Q9GJV1 ID Q9GJV1 PRELIMINARY; PRT; 9 AA.
AC Q9GJV1;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
DE ACTIN (FRAGMENT).
OS Lagenorhynchus acutus (Atlantic white-sided dolphin).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Cetartiodactyla; Cetacea; Odontoceti; Delphinidae;
OC Lagenorhynchus.
OX NCBI_TaxID=90246;
RN [1]
RP SEQUENCE FROM N.A.
RA Hare M.P., Cipriano F., Palumbi S.R.;
RT "Slow Evolution of Genetic Monophyly in Dolphins: Implications for
RT Speciation, Systematics and Conservation.";
RL Submitted (APR-1999) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF140825; AAF98678.1; -.
DR EMBL; AF140822; AAF98675.1; -.
DR EMBL; AF140823; AAF98676.1; -.
DR EMBL; AF140824; AAF98677.1; -.
FT NON_TER 1 1
FT NON_TER 9 9
SQ SEQUENCE 9 AA; 1049 MW; 1D0EF417740862C0 CRC64;

Query Match
Best Local Similarity 100.0%; Pred. No. 5.6e+05; Indels 0; Gaps 0;
Matches 2; Conservative 0; Mismatches 0;

QY 2 HQ 3
DB 8 HQ 9

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Query Match 23.6%; Score 13; DB 6; Length 9;
Best Local Similarity 100.0%; Pred. No. 5.6e+05;
Matches 2; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 2 HQ 3
II:
Db 8 HQ 9

RESULT 40
Q9T688
ID Q9T688 PRELIMINARY; PRT; 9 AA.
AC Q9T688;
DT 01-MAY-2000 (TReMBLrel. 13, Created)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Gecko gecko (Tokay gecko).
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Scleroglossa; Gekkota; Gekkonidae; Gekko.
OX NCBI_TaxID=36310;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99343618; PubMed=10413626;
RA Macey J.R., Wang Y., Ananjeva N.B., Larson A., Papenfuss T.J.;
RT "Variant patterns of fragmentation among gekkonid lizards of the
RT genus teratocincus produced by the Indian collision: A molecular
RT phylogenetic perspective and an area cladogram for central asia."
RL Mol. Phylogenet. Evol. 12:320-332(1999).
DR EMBL; AF114249; AAD51600.1; -.
KW Mitochondrion.
FT NON_TER 9
SQ SEQUENCE 9 AA; 1188 MW; 428CB9C9D36411A7 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 9;
Best Local Similarity 66.7%; Pred. No. 5.6e+05;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
II:
Db 6 FFS 8

RESULT 41
Q9T4P9
ID Q9T4P9 PRELIMINARY; PRT; 10 AA.
AC Q9T4P9;
DT 01-MAY-2000 (TReMBLrel. 13, Created)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Liolaemus darwini.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Tropidurinae; Liolaemus.
OX NCBI_TaxID=109408;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=SDSU3477, AND SDSU3472;
RA Schulte J.A. II, Macey J.R., Espinoza R.E., Larson A.;
RT "Phylogenetic Relationships in the Iguanid Lizard Genus Liolaemus:
RT Multiple Origins of Viviparous Reproduction and a Phylogenetic
RT Evaluation of Andean Vicariance."
RL Biol. J. Linn. Soc. Lond. 0:0-0(2000).
DR EMBL; AF099274; AAF18928.1; -.
DR EMBL; AF099272; AAF18922.1; -.
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
II:
Db 7 FFS 9

RESULT 42
Q9ZIV3
ID Q9ZIV3 PRELIMINARY; PRT; 10 AA.
AC Q9ZIV3;
DT 01-MAY-1999 (TReMBLrel. 10, Created)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Diposaurus dorsalis.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Iguaninae; Diposaurus.
OX NCBI_TaxID=51217;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
RT example using four subfamilies of the lizard family Iguanidae."
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049857; AAD02514.1; -.
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1275 MW; 1A3580C9D36411A0 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
II:
Db 7 FFS 9

RESULT 43
Q9ZIV0
ID Q9ZIV0 PRELIMINARY; PRT; 10 AA.
AC Q9ZIV0;
DT 01-MAY-1999 (TReMBLrel. 10, Created)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Petrosaurus thalassinus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae;
OX Petrosaurus.
OX NCBI_TaxID=81826;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
RT example using four subfamilies of the lizard family Iguanidae."
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049858; AAD02517.1; -.
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;

Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 44

Q92YU7 PRELIMINARY; PRT; 10 AA.
AC Q92YU7;
DT 01-MAY-1999 (TREMBlrel. 10, Created)
DT 01-MAY-1999 (TREMBlrel. 10, Last sequence update)
DT 01-MAY-1999 (TREMBlrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Sator angustus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae; Sator.
OX NCBI_TaxID=43619;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
example using four subfamilies of the lizard family Iguanidae.";
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049859; AAD02520.1; -
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 45

Q92YU4 PRELIMINARY; PRT; 10 AA.
AC Q92YU4;
DT 01-MAY-1999 (TREMBlrel. 10, Created)
DT 01-MAY-1999 (TREMBlrel. 10, Last sequence update)
DT 01-MAY-1999 (TREMBlrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Sceloporus graciosus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae;
OC Sceloporus.
OX NCBI_TaxID=43625;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
example using four subfamilies of the lizard family Iguanidae.";
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049860; AAD02523.1; -
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1365 MW; 129780C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 46

Q92YU1 PRELIMINARY; PRT; 10 AA.
AC Q92YU1;
DT 01-MAY-1999 (TREMBlrel. 10, Created)
DT 01-MAY-1999 (TREMBlrel. 10, Last sequence update)
DT 01-MAY-1999 (TREMBlrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Uma scoparia.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae; Uma.
OX NCBI_TaxID=81829;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
example using four subfamilies of the lizard family Iguanidae.";
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049861; AAD02526.1; -
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

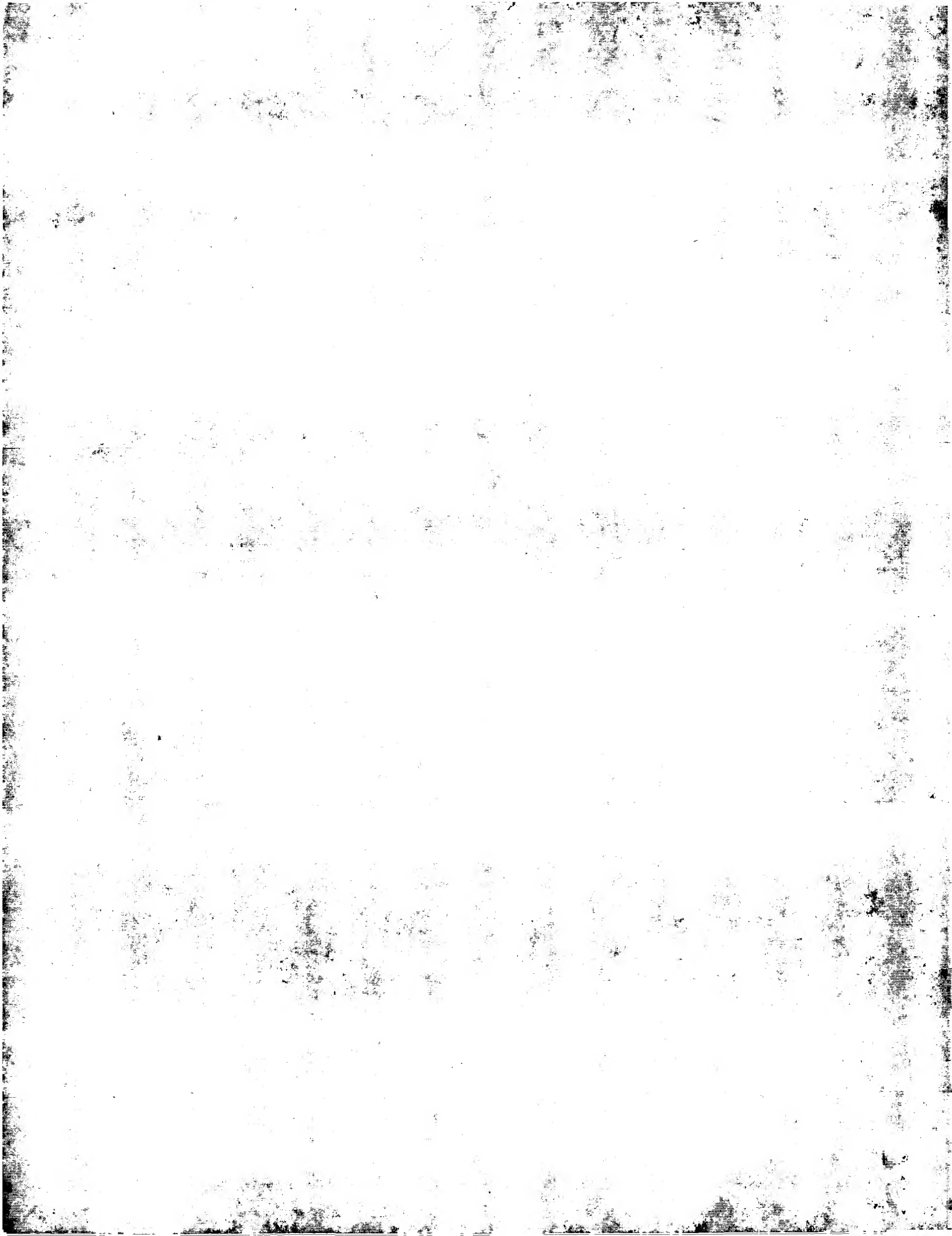
QY 7 FFA 9
||:
Db 7 FFS 9

RESULT 47

Q92YT8 PRELIMINARY; PRT; 10 AA.
AC Q92YT8;
DT 01-MAY-1999 (TREMBlrel. 10, Created)
DT 01-MAY-1999 (TREMBlrel. 10, Last sequence update)
DT 01-MAY-1999 (TREMBlrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Urosaurus graciosus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae;
OC Urosaurus.
OX NCBI_TaxID=43647;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT "Molecular tests of phylogenetic taxonomies: A general procedure and
example using four subfamilies of the lizard family Iguanidae.";
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049862; AAD02529.1; -
KW Mitochondrion.
FT NON_TER 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;
Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9
||:
Db 7 FFS 9



Db 7 FFS 9

RESULT 48

Q9ZYT5 PRELIMINARY; PRT; 10 AA.
AC Q9ZYT5
DT 01-MAY-1999 (TReMBLrel. 10, Created)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DT 01-MAY-1999 (TReMBLrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS utastansburiana.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Phrynosomatinae; Uta.
OX NCBI_TaxID=43653;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT Molecular tests of phylogenetic taxonomies: A general procedure and
RT example using four subfamilies of the lizard family Iguanidae.;
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049863; AAD02532.1; -.
KW Mitochondrion.
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;

Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9

||:

Db 7 FFS 9

RESULT 49

Q9ZYS9 PRELIMINARY; PRT; 10 AA.
AC Q9ZYS9
DT 01-MAY-1999 (TReMBLrel. 10, Created)
DT 01-MAY-1999 (TReMBLrel. 10, Last sequence update)
DT 01-MAY-1999 (TReMBLrel. 10, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Phymaturus somuncurensis.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Iguania; Iguanidae; Tropidurinae; Phymaturus.
OX NCBI_TaxID=81831;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99162288; PubMed=10051389;
RA Schulte J.A., Macey J.R., Larson A., Papenfuss T.J.;
RT Molecular tests of phylogenetic taxonomies: A general procedure and
RT example using four subfamilies of the lizard family Iguanidae.;
RL Mol. Phylogenet. Evol. 10:367-376(1998).
DR EMBL; AF049865; AAD02538.1; -.
KW Mitochondrion.
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1288 MW; 1A3580C9D3640440 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;

Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9

||:

Db 7 FFS 9

RESULT 50

Q9TG98 PRELIMINARY; PRT; 10 AA.
AC Q9TG98
DT 01-MAY-2000 (TReMBLrel. 13, Created)
DT 01-MAY-2000 (TReMBLrel. 13, Last sequence update)
DT 01-MAY-2000 (TReMBLrel. 13, Last annotation update)
DE CYTOCHROME C OXIDASE SUBUNIT I (FRAGMENT).
GN COI.
OS Shinisaurus crocodilurus.
OG Mitochondrion.
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Lepidosauria; Squamata; Scleroglossa; Anguilliforma; Shinisauridae;
OC Shinisaurus.
OX NCBI_TaxID=52224;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=99343613; PubMed=10413621;
RA Macey J.R., Schulte J.A. II, Larson A., Tuniyev B.S., Orlov N.,
RA Papenfuss T.J.;
RT Molecular phylogenetics, rRNA evolution, and historical biogeography
RT in anguillid lizards and related taxonomic families.;
RL Mol. Phylogenet. Evol. 12:250-272(1999).
DR EMBL; AF085604; AAD51502.1; -.
KW Mitochondrion.
FT NON_TER 10 10
SQ SEQUENCE 10 AA; 1290 MW; 1CEE80C9D36411A0 CRC64;

Query Match 23.6%; Score 13; DB 8; Length 10;
Best Local Similarity 66.7%; Pred. No. 3.5e+04;

Matches 2; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 7 FFA 9

||:

Db 7 FFS 9

Search completed: October 29, 2002, 09:38:29

Job time : 26 secs

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: October 29, 2002, 09:23:26 ; Search time 31 Seconds
(without alignments)
35,830 Million cell updates/sec

Title: US-09-724-842A-27
Perfect score: 55
Sequence: 1 HHQKLVFFAE 10

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 747574 seqs, 111073796 residues

Total number of hits satisfying chosen parameters: 747574

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 50 summaries

Database : A_Geneseq_032802.*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	55	100.0	10	22	Human APP derived
2	55	100.0	15	20	Beta-amyloid pepti
3	55	100.0	17	15	Beta-amyloid fragm
4	55	100.0	17	22	Amyloid beta-prote
5	55	100.0	17	22	Amyloid beta-prote
6	55	100.0	17	22	Beta-amyloid anti
7	55	100.0	18	21	Beta-amyloid precu
8	55	100.0	19	18	AEDANS-beta-amyloi
9	55	100.0	19	18	Trp-Beta-amyloid p
10	55	100.0	19	22	Human APP A-beta 1
11	55	100.0	19	22	Human amyloid beta

12	55	100.0	21	20	AA30941	Human secretase SE
13	55	100.0	24	15	AA52569	Alzheimer's disease
14	55	100.0	26	19	AA47229	Beta-amyloid pepti
15	55	100.0	26	20	AA33408	Human amyloidogeni
16	55	100.0	27	20	AA33409	Human amyloidogeni
17	55	100.0	28	8	AA70594	Sequence of Alzhei
18	55	100.0	28	10	AA90381	Synthetic A4 amylo
19	55	100.0	28	15	AA54702	Beta-amyloid fragm
20	55	100.0	28	15	AA60368	Beta-amyloid (1-28
21	55	100.0	28	16	AA64170	A4-O(1-28) a parti
22	55	100.0	28	16	AA64171	A4-P(1-28) a parti
23	55	100.0	28	16	AA64172	A4-B(1-28) a parti
24	55	100.0	28	16	AA64164	Generic beta amylo
25	55	100.0	28	17	AAW01413	Beta/A4-amyloid pe
26	55	100.0	28	20	AA39805	Beta-amyloid prote
27	55	100.0	28	20	AAW81467	Synthetic amyloid
28	55	100.0	28	22	AA91783	Amyloid beta-prote
29	55	100.0	28	22	AA91789	Amyloid beta-prote
30	55	100.0	28	22	AA91800	Amyloid beta-prote
31	55	100.0	28	22	AA91816	Amyloid beta-prote
32	55	100.0	28	22	AA91827	Amyloid beta-prote
33	55	100.0	28	22	AA43996	Human amyloid pept
34	55	100.0	28	22	AA35590	Human clone B(1-28
35	55	100.0	28	22	AA35591	Human clone D1N B(
36	55	100.0	28	22	AA35592	Human clone E3Q B(
37	55	100.0	28	22	AA35593	Human clone R5Q B(
38	55	100.0	28	22	AA35594	Human clone H6Q B(
39	55	100.0	28	22	AA35595	Human clone D7Q B(
40	55	100.0	28	22	AA35596	Human clone E11Q B
41	55	100.0	28	22	AA36201	Human clone D23Q B
42	55	100.0	28	22	AA36202	Human clone K28Q B
43	55	100.0	30	20	AAW81468	Synthetic amyloid
44	55	100.0	32	22	AA84430	Partial sequence o
45	55	100.0	33	20	AAW81469	Synthetic amyloid
46	55	100.0	35	19	AA47228	Beta-amyloid pepti
47	55	100.0	35	20	AAW89357	Beta-amyloid pepti
48	55	100.0	35	20	AAW89359	Beta-amyloid pepti
49	55	100.0	35	20	AAW89361	Beta-amyloid pepti
50	55	100.0	36	20	AAW81471	Synthetic amyloid

ALIGNMENTS

RESULT 1
ARB46225 ID AAB46225 standard; peptide; 10 AA.
XX
AC AAB46225;
XX
DT 04-APR-2001 (first entry)
XX
DE Human APP derived immunogenic peptide #21.
XX

XX Amyloid deposit; APP; Abeta; brain; human; clearing response; neurotropic;
KW Fc receptor mediated phagocytosis; immunogenic response; neuroprotective;
KW amyloid precursor protein; Alzheimer's disease.
XX
OS Homo sapiens.
XX
PN WO200072880-A2.
XX
PD 07-DEC-2000.
XX
PF 26-MAY-2000; 2000WO-US14810.
XX
PR 28-MAY-1999; 99US-0322289.
XX
PA (NEUR-) NEURALAB LTD.
XX
PI Schenk DB, Bard F, Vasquez NJ, Yednock T;
XX WPI; 2001-032104/04.
DR

Handwritten signature
Handwritten initials

XX Preventing or treating a disease associated with amyloid deposits,
 PT especially Alzheimer's disease, comprises administering amyloid
 PT specific antibody -
 XX
 PS Disclosure; Figure 19; 143pp; English.
 XX
 CC This invention describes a novel method of preventing or treating a
 CC disease associated with amyloid deposits of amyloid precursor protein
 CC (APP) Abeta fragments in the brain of a patient, which comprises
 CC administering to the patient: (a) an antibody that binds to Abeta, the
 CC antibody binds to an amyloid deposit and induces a clearing response (Fc
 CC receptor mediated phagocytosis) against it (b) a polypeptide containing
 CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
 CC that induces an immunogenic response against residues 1-3 to 7-11 of
 CC Abeta. The products of the invention have neurotropic and neuroprotective
 CC activity. The method is also useful for monitoring a course of treatment
 CC being administered to a patient e.g. active and passive immunization. The
 CC methods are useful for prophylactic and therapeutic treatment of
 CC Alzheimer's disease.
 XX
 SQ Sequence 10 AA;

Query Match 100.0%; Score 55; DB 22; Length 10;
 Best Local Similarity 100.0%; Pred. NO. 0.00014;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 DB 1 HHQKLVFFAE 10

RESULT 2
 AAW89358
 ID AAW89358 standard; peptide; 15 AA.
 XX
 AC AAW89358;

DT 02-MAR-1999 (first entry)

DE Beta-amyloid peptide derivative A-beta-11-25.

XX Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
 KW aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
 KW familial amyloid polynuropathy; bovine spongiform encephalopathy;
 KW Creutzfeldt-Jakob disease; bap.

XX Homo sapiens.
 OS Synthetic.

XX US5854204-A.

XX 29-DEC-1998.

XX 14-MAR-1996; 96US-0612785.

XX 14-MAR-1996; 96US-0612785.

XX 14-MAR-1995; 95US-0404831.

XX 07-JUN-1995; 95US-0475579.

XX 27-OCT-1995; 95US-0548998.

XX (PRAE-) PRAECIS PHARM INC.

XX Benjamin H, Chin J, Findeis MA, Garnick MB, Gelter ML;

PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;

PI Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;

XX WPI; 1999-094964/08.

XX New peptide(s) derived from beta-amyloid peptide that inhibit
 PT amyloid aggregation - and neurotoxicity, specifically for treatment
 PT and prevention of Alzheimer's disease

XX

PS Claim 6; Column 81-82; 52pp; English.

XX The present invention describes beta-amyloid peptide (bap) derivatives.
 CC The bap derivatives inhibit aggregation of amyloidogenic proteins and
 CC peptides, specifically bap, and their neurotoxicity, so are useful for
 CC treating and preventing any disease involving amyloidosis, specifically
 CC Alzheimer's disease but also Down's syndrome, familial amyloid
 CC polynuropathy or cardiomyopathy, bovine spongiform encephalopathy and
 CC Creutzfeldt-Jakob disease. The bap derivatives are also used to diagnose
 CC these diseases, in vitro or in vivo, by detecting binding of bap to
 CC labelled bap derivatives. Some bap derivatives inhibit bap aggregation
 CC even when bap is present in molar excess. The present sequence
 CC represents a bap derivative.

XX SQ Sequence 15 AA;

Query Match 100.0%; Score 55; DB 20; Length 15;
 Best Local Similarity 100.0%; Pred. NO. 0.00022;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 DB 3 HHQKLVFFAE 12

RESULT 3
 AAR54703
 ID AAR54703 standard; peptide; 17 AA.

XX AAR54703;

XX 15-DEC-1994 (first entry)

XX Beta-amyloid fragment (12-28).

XX Beta-amyloid protein; bap; Alzheimer's disease; diagnosis.

XX OS--Homo-sapiens.

XX WO9409364-A.

XX 28-APR-1994.

XX 13-OCT-1993; 93WO-US09772.

XX 13-OCT-1992; 92US-0959251.

XX (UYDU-) UNIV DUKE.

XX Strittmatter WJ;

XX WPI; 1994-151484/18.

XX Immobilised beta-amyloid protein or fragments - used in assays
 PT for obtaining prods for use in the diagnosis and treatment of
 PT disorders such as Alzheimer's disease.

XX Claim 5; Page 28; 49pp; English.

XX A construct comprising a beta-amyloid protein (BAP) or fragment (esp.
 CC the peptides given in AAR54702-03) immobilised on a solid support can be
 CC used to detect cpds. which bind to BAP. Binding of proteins in
 CC human cerebrospinal fluid proteins were shown to bind to beta-
 CC amyloid peptides 1-28 and 12-28. hydrophobic mimic peptide (12-28)
 CC was used as control.

XX SQ Sequence 17 AA;

Query Match 100.0%; Score 55; DB 15; Length 17;
 Best Local Similarity 100.0%; Pred. NO. 0.00025;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10

Db 2 HHQKLVFFAE 11

RESULT 4

AAB91774
ID AAB91774 standard; Peptide; 17 AA.

AC AAB91774;

XX 22-JUN-2001 (first entry)

DE Amyloid beta-protein fragment peptide SEQ ID NO:950.

KW Protection; endogenous therapeutic peptide; peptidase; conjugation;
KW blood component; modification; succinimidyl; maleimido group; amino;
KW hydroxyl; thiol; hormone; growth factor; neurotransmitter.

XX Homo sapiens.

OS Synthetic.

PN WO200069900-A2.

XX 23-NOV-2000. *13-2-00 11-29-00*

XX 17-MAY-2000; 2000WO-US13576.

XX 17-MAY-1999; 99US-0134406.

PR 10-SEP-1999; 99US-0153406.

PR 15-OCT-1999; 99US-0159783.

XX (CONJ-) CONJUCHEM INC.

XX Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;

XX WPI; 2001-112059/12.

XX Modifying and attaching therapeutic peptides to albumin prevents
PT peptidase degradation, useful for increasing length of in vivo activity
PT
PT
PS Disclosure; Page 504; 733pp; English.

XX The present invention describes a modified therapeutic peptide (I)
CC comprising a therapeutically active amino acid region (III) and a
CC reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
CC a less therapeutically active amino acid region (IV), which covalently
CC bonds with amino/hydroxyl/thiol groups on blood components to form a
CC peptidase stabilised therapeutic peptide composed of 3-50 amino acids.
CC (I) are useful for modifying therapeutic peptides e.g. hormones, growth
CC factors and neurotransmitters, to protect them from peptidase activity
CC in vivo for the treatment of various disorders. Endogenous therapeutic
CC peptides are not suitable as drug candidates as they require frequent
CC administration due to rapid degradation by peptidases in the body.
CC Modifying and attaching therapeutic peptides to albumin prevents or
CC reduces the action of peptidases to increase length of activity (half
CC life) and specificity as bonding to large molecules decreases
CC intracellular uptake and interference with physiological processes.
CC AAB90829 to AAB92441 represent peptides which can be used in the
CC exemplification of the present invention.

XX Sequence 17 AA;

Query Match 100.0%; Score 55; DB 22; Length 17;

Best Local Similarity 100.0%; Pred. No. 0.00025;

Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10

DB 2 HHQKLVFFAE 11

RESULT 5

AAB91807

ID AAB91807 standard; Peptide; 17 AA.

AC AAB91807;

XX 22-JUN-2001 (first entry)

DE Amyloid beta-protein fragment peptide SEQ ID NO:983.

KW Protection; endogenous therapeutic peptide; peptidase; conjugation;
KW blood component; modification; succinimidyl; maleimido group; amino;
KW hydroxyl; thiol; hormone; growth factor; neurotransmitter.

XX Homo sapiens.

OS Synthetic.

PN WO200069900-A2.

XX 23-NOV-2000.

XX 17-MAY-2000; 2000WO-US13576.

XX 17-MAY-1999; 99US-0134406.

PR 10-SEP-1999; 99US-0153406.

PR 15-OCT-1999; 99US-0159783.

XX (CONJ-) CONJUCHEM INC.

XX Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;

XX WPI; 2001-112059/12.

XX Modifying and attaching therapeutic peptides to albumin prevents
PT peptidase degradation, useful for increasing length of in vivo activity
PT
PT
PS Disclosure; Page 516; 733pp; English.

XX The present invention describes a modified therapeutic peptide (I)
CC comprising a therapeutically active amino acid region (III) and a
CC reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
CC a less therapeutically active amino acid region (IV), which covalently
CC bonds with amino/hydroxyl/thiol groups on blood components to form a
CC peptidase stabilised therapeutic peptide composed of 3-50 amino acids.
CC (I) are useful for modifying therapeutic peptides e.g. hormones, growth
CC factors and neurotransmitters, to protect them from peptidase activity
CC in vivo for the treatment of various disorders. Endogenous therapeutic
CC peptides are not suitable as drug candidates as they require frequent
CC administration due to rapid degradation by peptidases in the body.
CC Modifying and attaching therapeutic peptides to albumin prevents or
CC reduces the action of peptidases to increase length of activity (half
CC life) and specificity as bonding to large molecules decreases
CC intracellular uptake and interference with physiological processes.
CC AAB90829 to AAB92441 represent peptides which can be used in the
CC exemplification of the present invention.

XX Sequence 17 AA;

Query Match 100.0%; Score 55; DB 22; Length 17;

Best Local Similarity 100.0%; Pred. No. 0.00025;

Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10

DB 2 HHQKLVFFAE 11

RESULT 6

AAB48346

ID AAB48346 standard; peptide; 17 AA.

XX AAB48346;

XX

DT 20-APR-2001 (first entry)
 XX Beta-amyloid antigenic peptide (Abeta10-25).
 DE Beta-amyloid; nootropic; neuroprotective; vaccine; antibody; brain;
 XX amyloid plaque; Alzheimer's disease; antigen.
 KW Homo sapiens.
 OS
 XX Key Location/Qualifiers
 FH Modified-site 17 /note= "C-terminal amide"
 FT
 XX WO200077178-A1.
 PN 21-DEC-2000.
 XX 15-JUN-2000; 2000WO-US16551.
 PF 16-JUN-1999; 99US-0139408.
 XX (BOST-) BOSTON BIOMEDICAL RES INST.
 XX Raso V;
 PI WPI; 2001-112220/12.
 DR New antibodies which catalyze hydrolysis of beta-amyloid at a
 XX predetermined amide linkage, useful for e.g. sequestering or reducing
 PT free beta-amyloid in the bloodstream and brain and preventing formation
 PT of amyloid plaques -
 XX Example 1; Fig 3; 82pp; English.
 PS The invention relates to an antibody which catalyzes the hydrolysis of
 XX beta-amyloid at a predetermined amide linkage. The antibodies are useful
 CC for sequestering free beta-amyloid in the bloodstream of an animal,
 CC reducing beta-amyloid levels in the brain, preventing formation of
 CC amyloid plaques, and disaggregating amyloid plaques present in the brain,
 CC thus may be used in treating patients diagnosed with or at risk for
 CC Alzheimer's disease. The present sequence represents a beta-amyloid
 CC antigenic peptide made from the central region of beta-amyloid. The
 CC antigenic peptides were designed to be tested for suitability to
 CC antibody-mediated therapy.
 CC
 SQ Sequence 17 AA;
 Query Match 100.0%; Score 55; DB 22; Length 17;
 Best Local Similarity 100.0%; Pred. No. 0.00025;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 5 HHQKLVFFAE 14
 |||||
 RESULT 7
 AAB10963
 ID AAB10963 standard; protein; 18 AA.
 XX
 AC AAB10963;
 XX
 DT 07-FEB-2001 (first entry)
 DE Beta-amyloid precursor protein peptide fragment.
 XX APP; amyloid precursor protein; human; alpha-secretase; ADAM 10;
 KW disintegrin-metalloprotease; protease; nootropic; neuroprotective;
 KW gene therapy; Alzheimer's disease.
 XX Unidentified.
 OS
 XX DF19910108-A1.
 PN

XX 21-SEP-2000.
 XX 08-MAR-1999; 99DE-1010108.
 XX 08-MAR-1999; 99DE-1010108.
 XX (FAHR/) FAHRENHOLZ F.
 XX Fahrenholz F, Postina R;
 PI WPI; 2000-588391/56.
 DR
 XX Recombinant cells, for identifying alpha-secretase active agents and
 PT identifying risk factors associated with Alzheimer's disease, comprise
 PT amyloid precursor protein and alpha-secretase -
 XX Example 13; Page 12; 24pp; German.
 PS
 XX This invention describes a novel recombinant cell comprising recombinant
 CC nucleic acids encoding a region of human amyloid precursor protein
 CC containing an alpha-secretase cleavage site and a protease or a
 CC heterologous RNA coding for a substrate protein and a protease. The
 CC invention also describes a recombinant cell, characterized in that it
 CC contains recombinant nucleic acids comprising either: (a) a gene for a
 CC substrate protein (SP), which comprises a sequence region of 18 amino
 CC acids of the human amyloid precursor protein (APP) or a homologous
 CC protein, where the sequence region contains the alpha-secretase cleavage
 CC site at a reference of 6 residues at the N-terminal and 12 residues at
 CC the C-terminal; and (b) a gene for a protease protein (PP), that either
 CC comprises a proteolytically active necessary sequence region or a cow
 CC sequence region of the disintegrin metalloprotease ADAM 10 from a cow
 CC (Bos taurus), from a human or other mammal or a mutant of this, which
 CC shows the same enzymatic properties, where the genes are under the
 CC control of heterologous promoters; or a heterologous RNA coding for a SP
 CC and a PP. The products of the invention have nootropic and
 CC neuroprotective activity and can be used for gene therapy. The protease
 CC proteins of the invention are useful for proteolytic cleavage of
 CC substrate proteins, especially human amyloid precursor protein. Dominant
 CC negative forms of bovine, human or other mammalian
 CC disintegrin-metalloprotease ADAM 10 proteins and their coding sequences
 CC are useful for suppressing the alpha-secretase activity of a cell.
 CC Nucleic acid sequences encoding the proteases are useful for
 CC constructing vectors for gene therapy. The proteins and recombinant cells
 CC are useful for identifying secretases and pharmaceutical agents and to
 CC identify risk factors associated with Alzheimer's disease.
 CC
 SQ Sequence 18 AA;
 Query Match 100.0%; Score 55; DB 21; Length 18;
 Best Local Similarity 100.0%; Pred. No. 0.00027;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 3 HHQKLVFFAE 12
 |||||
 RESULT 8
 AAW18882
 ID AAW18882 standard; peptide; 19 AA.
 XX
 AC AAW18882;
 XX
 DT 08-DEC-1997 (first entry)
 DE AEDANS-beta-amyloid peptide fragment (9-25).
 XX beta-amyloid peptide; membrane protein; amyloid precursor protein;
 KW fibril assembly; in vitro; detection; fluorescence; amyloidosis disorder;
 KW Alzheimer's disease; multiple myeloma; rheumatoid arthritis; diabetes;
 KW prion disorder.
 XX

XX 26-MAY-2000; 2000WO-US14810.
 XX
 XX
 XX 28-MAY-1999; 99US-0322289.
 XX
 XX (NEUR-) NEURALAB LTD.
 XX
 XX
 XX Schenk DB, Bard F, Vasquez NJ, Yednock T;
 PI
 XX WPI; 2001-032104/04.
 DR
 XX Preventing or treating a disease associated with amyloid deposits,
 XX especially Alzheimer's disease, comprises administering amyloid
 PT specific antibody -
 PT
 XX Disclosure; Page 61; 143pp; English.
 PS
 XX This invention describes a novel method of preventing or treating a
 CC disease associated with amyloid deposits of amyloid precursor protein
 CC (APP) Abeta fragments in the brain of a patient, which comprises
 CC administering to the patient: (a) an antibody that binds to Abeta, the
 CC antibody binds to an amyloid deposit and induces a clearing response (Fc
 CC receptor mediated phagocytosis) against it (b) a polypeptide containing
 CC an N-terminal segment of at least residues 1-5 of Abeta; or (c) an agent
 CC that induces an immunogenic response against residues 1-3 to 7-11 of
 CC Abeta. The products of the invention have nootropic and neuroprotective
 CC activity. The method is also useful for monitoring a course of treatment
 CC being administered to a patient e.g. active and passive immunization. The
 CC methods are useful for prophylactic and therapeutic treatment of
 CC Alzheimer's disease.
 XX
 XX Sequence 19 AA;
 SQ

Query Match 100.0%; Score 55; DB 22; Length 19;
 Best Local Similarity 100.0%; Pred. No. 0.00029;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 Db 1 HHQKLVFFAE 10

RESULT 11
 AAB49097
 ID AAB49097 standard; peptide; 19 AA.
 XX
 XX AAB49097;
 XX
 XX 27-MAR-2001 (first entry)
 DT
 XX Human amyloid beta peptide (residues 13-28), SEQ ID NO:33.
 DE
 XX Amyloid disease; amyloid fibril deposition; amyloid plaque;
 KW immunogenic; antibody; vaccine; Alzheimer's disease;
 KW type 2 diabetes; reactive system amyloidosis;
 KW systemic senile amyloidosis; familial amyloid cardiomyopathy;
 KW transmissible spongiform encephalopathy; Creutzfeld-Jakob disease; Kuru;
 KW haemodialysis-associated beta-2-microglobulin deposition;
 KW amyloid beta peptide.
 XX
 XX Homo sapiens.
 OS
 XX WO200072876-A2.
 PN
 XX 07-DEC-2000.
 PD
 XX 01-JUN-2000; 2000WO-US15239.
 PF
 XX 01-JUN-1999; 99US-0137010.
 PR
 XX (NEUR-) NEURALAB LTD.
 XX
 XX Schenk DB;
 PI

XX WPI; 2001-070921/08.
 DR
 XX Pharmaceutical composition comprising immunogen against amyloid
 PT component such as fibril peptide or protein, or antibody against
 PT amyloid component useful for treating amyloid diseases or amyloidoses -
 XX
 XX Example IV; Page 74; 140pp; English.
 PS
 XX The invention relates to a novel pharmaceutical composition for
 CC preventing or treating a disease characterised by amyloid fibril
 CC deposits (amyloid plaques) in a patient. The pharmaceutical composition
 CC comprises an agent that will induce an immune response against an amyloid
 CC component, or an antibody or antibody fragment that binds to an amyloid
 CC component. The invention also relates to a method for determining
 CC the prognosis of a patient undergoing treatment for an amyloid disorder
 CC which involves measuring a patient serum amount of immunoreactivity
 CC against a selected amyloid component. A patient serum immunoreactivity
 CC of at least four times a base line serum immunoreactivity control level
 CC indicates a prognosis of improved status with respect to the disorder.
 CC The pharmaceutical compositions of the invention are useful for treating
 CC a wide variety of disorders characterised by amyloid fibril deposition in
 CC a patient. Such disorders include Alzheimer's disease characterised by
 CC amyloid beta peptide fibril deposits; type 2 diabetes characterised by
 CC amyloid beta peptide fibril deposits; type 2 diabetes characterised by
 CC amyloidosis associated with systemic inflammatory diseases (e.g.,
 CC rheumatoid arthritis, osteomyelitis, tuberculosis) characterised by AA
 CC fibrils derived from serum amyloid A protein (ApoSAA); systemic senile
 CC amyloidosis and familial amyloid cardiomyopathy characterised by ATTR
 CC fibrils derived from transthyretin (TTR); transmissible spongiform
 CC encephalopathies (e.g. Creutzfeld-Jakob disease, Kuru) characterised by
 CC prion protein deposits; and beta-2-microglobulin deposits which form as
 CC a result of long term haemodialysis treatment. The present sequence
 CC represents a human amyloid beta peptide which was conjugated to
 CC sheep anti-mouse IgG in an exemplification of the invention.
 XX
 XX Sequence 19 AA;
 SQ

Query Match 100.0%; Score 55; DB 22; Length 19;
 Best Local Similarity 100.0%; Pred. No. 0.00029;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 Db 1 HHQKLVFFAE 10

RESULT 12
 AAY30941
 ID AAY30941 standard; peptide; 21 AA.
 XX
 XX AAY30941;
 XX
 XX 19-OCT-1999 (first entry)
 DT
 XX Human secretase SEC-alpha1 peptide fragment.
 DE
 XX Secretase; hyperforin; treatment; Alzheimer's disease; purification;
 KW adhyperforin; St. John's Wort; storage stable; pharmaceutical;
 KW symptom; SEC-alpha1; human.
 KW
 XX Homo sapiens.
 OS
 XX WO9941220-A1.
 PN
 XX 19-AUG-1999.
 PD
 XX 04-FEB-1999; 99WO-EP00737.
 PF
 XX 13-FEB-1998; 98DE-1005947.
 PR
 XX (SCHW-) SCHWABE GMBH & CO WILLMAR.
 PA
 XX
 XX

PI Chatterjee SS, Erdelmeier C, Klessing K, Marne D;
 PI Schaechtele C;
 DR WPI; 1999-508609/42.
 XX
 XX Hyperforin and adhyperforin isolated from St. John's Wort for
 PT treatment of Alzheimers
 PT
 XX
 PS Example 34; Fig 1; 41pp; German.
 CC
 CC This invention describes novel hyperforin and adhyperforin salts of
 CC formula (I): (A-)m (B)p+, where m = 1-3; (A-) = an anion of formula (II);
 CC n = 0-1; (B)p+ = an alkali metal ion or an ammonium ion of a salt-forming
 CC nitrogen base of formula (III); R1-R3 = H, an optionally branched alkyl,
 CC cycloalkyl, bicycloalkyl, tricycloalkyl, alkenyl, alkynyl,
 CC heterocycloalkyl, aryl, heteroaryl, arylalkyl or a heteroarylalkyl group,
 CC all optionally substituted with one or more hydroxy, alkoxy, aryloxy,
 CC alkanoyl, aroyl, carboxy, alkoxycarbonyl, ureido, amidino, guanidino,
 CC cyano, azido, mercapto, alkylthio, alkylsulphoxy, alkylsulphonyl,
 CC alkylsulphenyl, aminosulphonyl, fluoro, chloro, bromo, iodo, alkyl or
 CC perfluoroalkyl; R1+R2 = together with an N-atom form, together with a
 CC N-atom an azetidin-, pyrrolidin-, pyrrolin-, piperidin-, piperazin-,
 CC homopiperazin-, morpholin-, thiomorpholin-, pyridin-, di- or
 CC tetra-hydropyridin-, pyrimidin-, pyrazin-, azeplin-, dihydroazepin-,
 CC oxazepin-, diazepin-, imidazol-, pyrazol-, oxazol- or thiazol-ring,
 CC optionally with aliphatic, heteroaliphatic, aromatic or heteroaromatic
 CC rings or substituted with hydroxy, alkoxy, aryloxy, alkanoyl, aroyl,
 CC carboxy, alkoxycarbonyl, ureido, amidino, guanidino, cyano, azido,
 CC mercapto, alkylthio, alkylsulphoxy, alkylsulphonyl, alkylsulphenyl,
 CC aminosulphonyl, fluoro, chloro, bromo, iodo, alkyl or perfluoroalkyl;
 CC R4 = H, or an optionally branched alkyl group. The preparation is used to
 CC purify the hyperforin and/or adhyperforin content in St. John's Wort
 CC extracts. The obtained salts are storage stable and can be used in
 CC pharmaceutical compositions for the treatment of Alzheimer's disease and
 CC its symptoms. This sequence represents a fragment of the human secretase
 CC Sec-alpha1 protein which is used to illustrate the method of the
 CC invention.
 XX
 XX
 SQ Sequence 21 AA;

Query Match 100.0%; Score 55; DB 20; Length 21;
 Best Local Similarity 100.0%; Pred. NO. 0.00032;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 Db 8 HHQKLVFFAE 17

RESULT 13

AAR52569
 ID AAR52569 standard; peptide; 24 AA.

XX
 AC AAR52569;

DT 16-DEC-1994 (first entry)

XX Alzheimer's disease related immunogen.

XX Alzheimer's disease; senile dementia; immunogen.

XX Synthetic.

XX JP06009693-A.

XX 18-JAN-1994.

XX 23-JAN-1992; 92JP-0031341.

XX 23-JAN-1992; 92JP-0031341.

XX (EIKE) EIKEN KAGAKU KK.

XX

DR
 XX

PT Alzheimer's disease related protein isolated from serum of
 PT patient - useful in diagnosis
 XX

PS Claim 1; Page 2; 8pp; Japanese.

XX A monoclonal antibody raised against the synthetic peptide AAR52569 as
 CC immunogen reacts with a new Alzheimer's disease related protein. The
 CC novel protein has a mol.wt. of 20kD (by SDS-PAGE), isoelectric point
 CC of ca. 5-7 and is abundant in serum of AD patients.

XX Sequence 24 AA;

Query Match 100.0%; Score 55; DB 15; Length 24;
 Best Local Similarity 100.0%; Pred. No. 0.00037;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 |||||
 Db 13 HHQKLVFFAE 22

RESULT 14

AAR52569
 ID AAR52569 standard; peptide; 26 AA.

XX
 AC AAR52569;

DT 22-MAY-1998 (first entry)

XX Beta-amyloid peptide residues 10-35.

XX Screening assay; beta-amyloid peptide; treatment;
 KW amyloidosis disease; Alzheimer's disease.

XX Homo sapiens.

XX US5721106-A.

XX 24-FEB-1998.

PF 12-SEP-1994; 94US-0304585.

PR 12-SEP-1994; 94US-0304585.

PR 13-AUG-1991; 91US-0744767.

XX (HARD) HARVARD COLLEGE.

PA (MINU) UNIV MINNESOTA.

XX Maggio JE, Mantyh PW;

XX WPI; 1998-168404/15.

XX New in vitro screening assay for Alzheimer's disease drugs -
 PT comprises assessing binding of labelled beta-amyloid peptide to silk
 PT sample

PS Claim 8; Columns 31-32; 36pp; English.

XX The present sequence was used in the development of a novel in
 CC vitro screening assay for agents capable of affecting the
 CC deposition of beta-amyloid peptide (BAP) on tissue. The method
 CC comprises contacting a silk sample with labelled BAP, optionally
 CC in the presence of a test agent, detecting the amount of label
 CC bound to the silk and assessing the effect of the agent on the
 CC deposition of BAP. Agents that inhibit binding of BAP to silk are
 CC potentially useful for treating amyloidosis diseases, especially
 CC Alzheimer's disease.

XX Sequence 26 AA;

Query Match 100.0%; Score 55; DB 19; Length 26;

Best Local Similarity 100.0%; Pred. No. 0.0004; Mismatches 0; Indels 0; Gaps 0;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 4 HHQKLVFFAE 13

RESULT 15
AAY33408
ID AAY33408 standard; peptide; 26 AA.
XX
AC AAY33408;
XX
DT 03-DEC-1999 (first entry)
XX
DE Human amyloidogenic A-beta peptide 2.
XX
KW Amyloidogenic; beta-amyloid; A-beta peptide; human; inhibitor;
KW fibrillogenesis; amyloid plaque; amyloidosis; Alzheimer's disease;
KW Down's Syndrome.
XX
OS Homo sapiens.
XX
PN WO9941279-A2.
XX
PD 19-AUG-1999.
XX
PF 12-FEB-1999; 99WO-US03231.
XX
PR 13-FEB-1998; 98US-0074658.
XX
PA (ARCH-) ARCH DEV CORP.
XX
PI Lynn DG, Meredith SC, Burkoth TS;
XX
PT Inhibiting amyloid plaque formation in humans suffering from
PT amyloidosis, Alzheimer's disease or Down's Syndrome -
XX
PS Claim 22; Page 140; 141pp; English.
XX
CC This invention describes a novel method for inhibiting amyloid
CC fibrillogenesis which comprises contacting tissue with a composition
CC comprising an amyloidogenic peptide, beta-amyloid, that has been blocked
CC at an end terminal or a side chain, by conjugation to polyethylene
CC glycol, by conjugation to a second compound and a pharmaceutically
CC acceptable buffer, solvent or diluent. The methods are used to inhibit
CC amyloid plaque formation in humans suffering from amyloidosis,
CC Alzheimer's disease or Down's Syndrome. This sequence represents a
CC fragment of the beta-amyloid peptide described in the method of the
XX invention.
XX
SQ Sequence 26 AA;
Query Match 100.0%; Score 55; DB 20; Length 26;
Best Local Similarity 100.0%; Pred. No. 0.0004;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 4 HHQKLVFFAE 13

RESULT 16
AAY33409
ID AAY33409 standard; peptide; 27 AA.
XX
AC AAY33409;
XX
DT 03-DEC-1999 (first entry)
XX

DE Human amyloidogenic A-beta peptide C-terminal fragment.
XX
KW Amyloidogenic; beta-amyloid; A-beta peptide; human; inhibitor;
KW fibrillogenesis; amyloid plaque; amyloidosis; Alzheimer's disease;
KW Down's Syndrome.
XX
OS Homo sapiens.
XX
PN WO9941279-A2.
XX
PD 19-AUG-1999.
XX
PF 12-FEB-1999; 99WO-US03231.
XX
PR 13-FEB-1998; 98US-0074658.
XX
PA (ARCH-) ARCH DEV CORP.
XX
PI Lynn DG, Meredith SC, Burkoth TS;
XX
PT Inhibiting amyloid plaque formation in humans suffering from
PT amyloidosis, Alzheimer's disease or Down's Syndrome -
XX
PS Disclosure; Page 141; 141pp; English.
XX
CC This invention describes a novel method for inhibiting amyloid
CC fibrillogenesis which comprises contacting tissue with a composition
CC comprising an amyloidogenic peptide, beta-amyloid, that has been blocked
CC at an end terminal or a side chain, by conjugation to polyethylene
CC glycol, by conjugation to a second compound and a pharmaceutically
CC acceptable buffer, solvent or diluent. The methods are used to inhibit
CC amyloid plaque formation in humans suffering from amyloidosis,
CC Alzheimer's disease or Down's Syndrome. This sequence represents the
CC C-terminal fragment of a PEG-derivatized beta-amyloid peptide described
XX in the method of the invention.
XX
SQ Sequence 27 AA;
Query Match 100.0%; Score 55; DB 20; Length 27;
Best Local Similarity 100.0%; Pred. No. 0.00042;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 HHQKLVFFAE 10
Db 5 HHQKLVFFAE 14

RESULT 17
AAP70594
ID AAP70594 standard; peptide; 28 AA.
XX
AC AAP70594;
XX
DT 15-APR-1991 (first entry)
XX
DE Sequence of Alzheimer's amyloid polypeptide (AAP).
XX
KW Diagnosis; immunologic assay.
XX
OS Homo sapiens.
XX
PN US4666829-A.
XX
PD 19-MAY-1987.
XX
PF 15-MAY-1985; 85US-0734660.
XX
PR 15-MAY-1985; 85US-0734660.
XX
PA (REGC) UNIV OF CALIFORNIA.
XX

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PI  Glenner GG, Wong CW;
XX  WPI; 1987-157148/22.
XX
XX  Alzheimer's amyloid polypeptide - used for obtaining antibodies
XX  and nucleotide probes for diagnosis of Alzheimer's disease
XX
XX  Claim 1; column 11; 8pp; English.
XX
XX  Brains obtd. from patients suspected of having Alzheimer's disease
XX  and exhibiting extensive cerebrovascular amyloidosis were used for
XX  AAP isolation. The AAP can be used to obtain antibodies which can
XX  be used as reagents (claimed) in a blood or tissue immunologic
XX  assay for the disease. It can also be used to develop a probe
XX  (claimed) which can be used in a diagnostic test (claimed).
XX
XX  Sequence 28 AA;
SQ
Query Match      100.0%; Score 55; DB 8; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY  1 HHQKLVFFAE 10
Db  13 HHQKLVFFAE 22

RESULT 18
AAP90381
ID  AAP90381 standard; protein; 28 AA.
XX
XX  AAP90381;
XX
XX  01-NOV-1989 (first entry)
XX
XX  Synthetic A4 amyloid peptide.
XX
XX  Synthetic; A4 amyloid polypeptide; Alzheimer's disease;
XX  immunoassays; antibodies.
XX
XX  Synthetic.
XX
XX  W08906242-A.
XX
XX  13-JUL-1989.
XX
XX  11-OCT-1988; 88WO-US03590.
XX
XX  08-OCT-1987; 87US-0105751.
XX
XX  (MCLE ) MCLEAN HOSPITAL CORP; (UYRO) UNIVERSITY OF ROCHESTER.
XX
XX  Majocha R, Marotta CA, Zain S;
XX  WPI; 1989-220551/30.
XX
XX  Antibodies to A4 amyloid polypeptide
XX  - used in immunoassays and for imaging of A4 amyloid
XX  in Alzheimer's diseased patients.
XX
XX  Claim 1; page 27; 30pp; English.
XX
XX  Synthetic A4 amyloid polypeptide (see also AAP90382, AAP90383).
XX  used as immunogen, (un)coupled, or to produce antibodies. Used in
XX  immunoassays and for imaging of A4 amyloid in Alzheimer's disease.
XX
XX  Sequence 28 AA;
SQ
Query Match      100.0%; Score 55; DB 10; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY  1 HHQKLVFFAE 10

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Db  13 HHQKLVFFAE 22

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RESULT 19
AAR54702
ID  AAR54702 standard; peptide; 28 AA.
XX
XX  AAR54702;
XX
XX  15-DEC-1994 (first entry)
XX
XX  Beta-amyloid fragment (1-28).
XX
XX  Beta-amyloid protein; BAP; Alzheimer's disease; diagnosis.
XX
XX  Homo sapiens.
XX
XX  W09409364-A.
XX
XX  28-APR-1994.
XX
XX  13-OCT-1993; 93WO-US09772.
XX
XX  13-OCT-1992; 92US-0959251.
XX
XX  (UYDU-) UNIV DUKE.
XX
XX  Strittmatter WJ;
XX
XX  WPI; 1994-151484/18.
XX
XX  Immobilised beta-amyloid protein or fragments - used in assays
XX  for obtaining prods for use in the diagnosis and treatment of
XX  disorders such as Alzheimer's disease.
XX
XX  Claim 4; Page 28; 49pp; English.
XX
XX  A construct comprising a beta-amyloid protein (BAP) or fragment (esp.
XX  the peptides given in AAR54702-03) immobilised on a solid support can be
XX  used to detect cpds. which bind to BAP. Binding of proteins in
XX  human cerebrospinal fluid proteins were shown to bind to beta-
XX  amyloid peptides 1-28 and 12-28. Hydropathic mimic peptide (12-28)
XX  was used as control.
XX
XX  Sequence 28 AA;
SQ
Query Match      100.0%; Score 55; DB 15; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY  1 HHQKLVFFAE 10
Db  13 HHQKLVFFAE 22

```

```

RESULT 20
AAR60368
ID  AAR60368 standard; peptide; 28 AA.
XX
XX  AAR60368;
XX
XX  15-MAR-1995 (first entry)
XX
XX  Beta-amyloid (1-28).
XX
XX  Amyloid precursor protein; APP; Alzheimer's disease; beta-amyloid;
XX  anti-beta-amyloid antibody; diagnosis; immunogen; antigen; epitope.
XX
XX  Homo sapiens.
XX
XX  W09417197-A.
XX

```



```
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 23
AAR64172
ID AAR64172 standard; peptide; 28 AA.
XX
AC AAR64172;
XX
DT 03-AUG-1995 (first entry)
XX
DE A4-B(1-28) a partial beta amyloid peptide.
XX
KW beta amyloid protein; mutant; variant; detection; amyloid deposition;
KW diagnosis; amyloidosis associated disease; Alzheimer's disease;
KW Down's syndrome; A4-B(1-28).
XX
OS Synthetic.
XX
PN WO9428412-A.
XX
PD 08-DEC-1994.
XX
PF 27-MAY-1994; 94WO-US05809.
XX
PR 28-MAY-1993; 93US-0069010.
XX
PA (MIRI-) MIRIAM HOSPITAL.
XX
PI Majocha RE, Marotta CA;
XX
WPI; 1995-023013/03.
XX
PT Amyloid binding composition comprising labelled amyloid protein
PT and carrier - useful for in vivo imaging of amyloid deposits, for
PT diagnosing Alzheimer's disease and Down's Syndrome.
XX
PS Example 3; Page 23; 58pp; English.
XX
CC AAR64172, the A4-B(1-28) polypeptide is deriv. from vascular amyloid of
CC the AD (Alzheimer's disease) brain and a Down Syndrome brain. Three of
CC the 28 amino acids are different from the A4-O(1-28) peptide shown in
CC AAR64170. A4-O has strong aggregation properties, and binds to itself
CC strongly. It is used to obtain and select beta amyloid proteins that can
CC be used for in vivo imaging of amyloid deposits and hence diagnosis of
CC an amyloidosis-associated disease, such as AD or Down's syndrome.
CC AAR64165 shows the generic sequence of the amyloid protein for generation
CC of variants.
XX
SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 16; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 24
AAR64164
ID AAR64164 standard; peptide; 28 AA.
XX
AC AAR64164;
XX
DT 02-AUG-1995 (first entry)
XX
DE Generic beta amyloid protein variant.
```

```
XX generic sequence; beta amyloid protein; mutant; variant; detection;
KW amyloid deposition; diagnosis; amyloidosis associated disease;
KW Alzheimer's disease; Down's syndrome.
XX
OS Synthetic.
XX
XX Key Location/Qualifiers
FT Misc-difference 11 /note= "Glu or Gln"
FT Misc-difference 27 /note= "Ser or Asn"
FT Misc-difference 28 /note= "Ala or Lys"
XX
PN WO9428412-A.
XX
PD 08-DEC-1994.
XX
PF 27-MAY-1994; 94WO-US05809.
XX
PR 28-MAY-1993; 93US-0069010.
XX
PA (MIRI-) MIRIAM HOSPITAL.
XX
PI Majocha RE, Marotta CA;
XX
WPI; 1995-023013/03.
XX
PT Amyloid binding composition comprising labelled amyloid protein
PT and carrier - useful for in vivo imaging of amyloid deposits, for
PT diagnosing Alzheimer's disease and Down's Syndrome.
XX
PS Claim 3; Page 42; 58pp; English.
XX
CC AAR64164 shows the generic amino acid sequence of a variant beta amyloid
CC protein. The protein binds amyloid and is useful for in vivo imaging of
CC amyloid deposits and hence diagnosis of an amyloidosis-associated
CC disease, such as Alzheimer's disease or Down's syndrome. AAR64165-69
CC show specific variants generated from this generic sequence with addition
CC amino acids.
XX
SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 16; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
Db 13 HHQKLVFFAE 22

RESULT 25
AAR64163
ID AAR64163 standard; Protein; 28 AA.
XX
AC AAR64163;
XX
DT 20-JAN-1997 (first entry)
XX
DE Beta/A4-amyloid peptide residues 1-28.
XX
KW Beta/A4-amyloid peptide; tissue plasminogen activator;
KW Alzheimer's disease; stimulation; investigation; pathogenesis;
KW hereditary cerebral haemorrhage with amyloidosis-Dutch type;
KW control; cerebral amyloid angiopathy; cerebral; haemorrhage;
XX
OS Homo sapiens.
XX
PN WO9615799-A1.
XX
```

PD 30-MAY-1996.
 XX
 PF 22-NOV-1995; 95WO-US15007.
 XX
 PR 22-NOV-1994; 94US-0347144.
 XX
 PA (RUTF) UNIV RUTGERS STATE NEW JERSEY.
 XX
 PI Anderson S;
 XX
 DR WPI; 1996-268332/27.
 XX
 PT Use of agents which bind beta-amyloid peptide - for diagnosis,
 PT prevention and treatment of vascular damage caused by amyloid
 PT deposits, partic. in haemorrhaging and Alzheimer's disease
 XX
 PS Example 1; Fig 1; 52pp; English.
 XX
 CC To investigate the effects of beta-amyloid peptide (BAP) on
 CC tissue plasminogen activator (t-PA) 3 synthetic peptides were used.
 CC One peptide contained 42 amino acids and corresp. to the full
 CC length BAP (AAR95248). The other 2 peptides (AAR95249 and 50) contained
 CC the 28 N-terminal residues of the BAP found in Alzheimer's disease
 CC and hereditary cerebral haemorrhage with amyloidosis-Dutch type
 CC (HCHWA-D), respectively. In an assay to determine the effect of
 CC the peptides on t-PA activation, each peptide (AAR95248, 49 and 50)
 CC gave 1st order rate constant of activation (k(app)) values of
 CC 13.4, 13.9 and 14.5, respectively, compared to 1.7 and 7.8 for nil
 CC and fibrinogen controls. The results demonstrate that the BAP are
 CC able to stimulate t-PA activity in vitro, which is significant in
 CC that it provides a means for investigating and controlling the
 CC pathogenesis of Alzheimer's disease, HCHWA-D and cerebral amyloid
 CC angioathy related cerebral haemorrhage.
 XX
 SQ Sequence 28 AA;
 Query Match 100.0%; Score 55; DB 17; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Qy 1 HHQKLVFFAE 10
 Db 13 HHQKLVFFAE 22
 RESULT 26
 AAY39805
 ID AAY39805 standard; peptide; 28 AA.
 XX
 AC AAY39805;
 XX
 DT 29-NOV-1999 (first entry)
 XX
 DE Beta-amyloid protein, Beta/A4 amyloid (1-28).
 XX
 KW Beta-amyloid protein; Alzheimer's disease; amyloidosis; joint swelling;
 KW long-standing inflammation; malignancy; Familial Mediterranean Fever;
 KW multiple myeloma; plasma cell dyscrasia; long-term haemodialysis; kuru;
 KW carpal tunnel syndrome; multiple spontaneous fracture; radiolucency;
 KW endocrine tumour; medullary carcinoma; Down's syndrome; scrapie;
 KW Creutzfeldt-Jakob disease; Gerstmann Strausler Syndrome;
 KW subacute spongiform encephalopathy; therapy.
 XX
 OS Homo sapiens.
 XX
 PN US5958883-A.
 XX
 PD 28-SEP-1999.
 XX
 PF 05-JUN-1995; 95US-0461216.
 XX
 PR 23-OCT-1992; 92US-0969734.
 PR 23-SEP-1992; 92US-0950417.

XX (UNIW) UNIV WASHINGTON.
 XX
 PI Snow AD;
 XX
 XX WPI; 1999-561062/47.
 DR
 XX Peptides of 6-8 amino acids useful for treating or preventing
 PT amyloidosis -
 PT
 XX Disclosure; Column 67-68; 83pp; English.
 PS
 XX This sequence represents a fragment of the beta-amyloid protein. The
 CC invention relates to a method for treating or preventing a form of
 CC amyloidosis, including Alzheimer's disease using this sequence. The
 CC compositions may be useful for treating or preventing the amyloidosis
 CC associated with long-standing inflammation, various forms of malignancy
 CC (including B-cell type malignancies), Familial Mediterranean Fever,
 CC multiple myeloma, plasma cell dyscrasias, long-term haemodialysis, carpal
 CC tunnel syndrome, joint swelling, multiple spontaneous fractures,
 CC radiolucency in the wrist and hip, endocrine tumours, medullary carcinoma
 CC of the thyroid, diabetes, Alzheimer's disease, Down's syndrome, scrapie
 CC Creutzfeldt-Jakob disease, Gerstmann Strausler Syndrome, kuru, scrapie
 CC and other subacute spongiform encephalopathies.
 XX
 SQ Sequence 28 AA;
 Query Match 100.0%; Score 55; DB 20; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Qy 1 HHQKLVFFAE 10
 Db 13 HHQKLVFFAE 22
 RESULT 27
 AAW81467
 ID AAW81467 standard; peptide; 28 AA.
 XX
 AC AAW81467;
 XX
 DT 28-JAN-1999 (first entry)
 XX
 DE Synthetic amyloid beta (Abeta) peptide 2 (residues 1-28).
 XX
 KW Amyloid beta; Abeta; deoxygenated solvent; evaporative deposition;
 KW research; neurotoxicity; free-radical; glutamine synthetase.
 XX
 OS Synthetic.
 XX
 PN US5840838-A.
 XX
 PD 24-NOV-1998.
 XX
 PF 29-FEB-1996; 96US-0609090.
 XX
 PR 29-FEB-1996; 96US-0609090.
 XX
 PA (KENT) UNIV KENTUCKY RES FOUND.
 XX
 PI Aksenov M, Butterfield DA, Carney JM, Hensley K;
 XX
 DR WPI; 1999-034120/03.
 XX
 PT Process for treating synthetic amyloid beta peptides - by organic
 PT solvent treatment, useful for studying neurotoxicity
 XX
 PS Claim 5; Columns 9-10; 14pp; English.
 XX
 CC Sequences AAW81466 to AAW81476 represent synthetic amyloid beta (Abeta)
 CC peptides. The invention provides a process for treating a synthetic
 CC Abeta peptide that comprises dissolving the peptide in a deoxygenated

CC solvent selected from trifluoroethanol, hexafluorocyclohexane, dimethyl
 CC sulphoxide, morpholinopropanesulphonic acid, dimethylformamide and
 CC acetonitrile to a concentration of 0.01-10 mg/ml, incubating the
 CC solution at 20-65 deg. C for 0.5-4 hour, and removing the solvent by
 CC /evaporative deposition'' in 5-10 minutes. Synthetic amyloid beta
 CC peptides are useful as research tools for studying neurotoxicity
 CC resulting from Abeta peptide -enhanced free-radical production. The
 CC treatment increases the activity of the synthetic Abeta peptides in tests
 CC to determine free-radical generating capacity and glutamine synthetase
 CC inactivation..
 XX
 SQ Sequence 28 AA;

Query Match 100.0%; Score 55; DB 20; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 DB 13 HHQKLVFFAE 22

RESULT 28

AAB911783
 ID AAB91783 standard; Peptide; 28 AA.

AC AAB91783;

DT 22-JUN-2001 (first entry)

DE Amyloid beta-protein fragment peptide SEQ ID NO:959.

Protection; endogenous therapeutic peptide; peptidase; conjugation;
 blood component; modification; succinimidyl; maleimido group; amino;
 hydroxyl; thiol; hormone; growth factor; neurotransmitter.

OS Homo sapiens.

OS Synthetic.

PN WO200069900-A2.

PD 23-NOV-2000.

PF 17-MAY-2000; 2000WO-US13576.

PR 17-MAY-1999; 99US-0134406.

PR 10-SEP-1999; 99US-0153406.

PR 15-OCT-1999; 99US-0159783.

(CONJ-) CONJUCHEM INC.

PI Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;

WPI; 2001-112059/12.

Modifying and attaching therapeutic peptides to albumin prevents
 peptidase degradation, useful for increasing length of in vivo activity

Disclosure; Page 507; 733pp; English.

The present invention describes a modified therapeutic peptide (I)
 comprising a therapeutically active amino acid region (III) and a
 reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
 a less therapeutically active amino acid region (IV), which covalently
 bonds with amino/hydroxyl/thiol groups on blood components to form a
 peptidase stabilised therapeutic peptide composed of 3-50 amino acids.
 (I) are useful for modifying therapeutic peptides e.g. hormones, growth
 factors and neurotransmitters, to protect them from peptidase activity
 in vivo for the treatment of various disorders. Endogenous therapeutic
 peptides are not suitable as drug candidates as they require frequent
 administration due to rapid degradation by peptidases in the body.
 Modifying and attaching therapeutic peptides to albumin prevents or

CC reduces the action of peptidases to increase length of activity (half
 CC life) and specificity as bonding to large molecules decreases
 CC intracellular uptake and interference with physiological processes.
 CC AAB90829 to AAB92441 represent peptides which can be used in the
 CC exemplification of the present invention.
 XX
 SQ Sequence 28 AA;

Query Match 100.0%; Score 55; DB 22; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVFFAE 10
 DB 13 HHQKLVFFAE 22

RESULT 29

AAB91789
 ID AAB91789 standard; Peptide; 28 AA.

AC AAB91789;

DT 22-JUN-2001 (first entry)

DE Amyloid beta-protein fragment peptide SEQ ID NO:965.

Protection; endogenous therapeutic peptide; peptidase; conjugation;
 blood component; modification; succinimidyl; maleimido group; amino;
 hydroxyl; thiol; hormone; growth factor; neurotransmitter.

OS Homo sapiens.

OS Synthetic.

PN WO200069900-A2.

XX 23-NOV-2000.

XX 17-MAY-2000; 2000WO-US13576.

XX 17-MAY-1999; 99US-0134406.

PR 10-SEP-1999; 99US-0153406.

PR 15-OCT-1999; 99US-0159783.

(CONJ-) CONJUCHEM INC.

PI Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;

WPI; 2001-112059/12.

Modifying and attaching therapeutic peptides to albumin prevents
 peptidase degradation, useful for increasing length of in vivo activity

Disclosure; Page 509; 733pp; English.

The present invention describes a modified therapeutic peptide (I)
 comprising a therapeutically active amino acid region (III) and a
 reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
 a less therapeutically active amino acid region (IV), which covalently
 bonds with amino/hydroxyl/thiol groups on blood components to form a
 peptidase stabilised therapeutic peptide composed of 3-50 amino acids.
 (I) are useful for modifying therapeutic peptides e.g. hormones, growth
 factors and neurotransmitters, to protect them from peptidase activity
 in vivo for the treatment of various disorders. Endogenous therapeutic
 peptides are not suitable as drug candidates as they require frequent
 administration due to rapid degradation by peptidases in the body.
 Modifying and attaching therapeutic peptides to albumin prevents or
 reduces the action of peptidases to increase length of activity (half
 CC life) and specificity as bonding to large molecules decreases
 CC intracellular uptake and interference with physiological processes.
 CC AAB90829 to AAB92441 represent peptides which can be used in the
 CC exemplification of the present invention.

```

XX SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
DB 13 HHQKLVFFAE 22

RESULT 30
AAB91800
ID AAB91800 standard; Peptide; 28 AA.
AC AAB91800;
XX
XX
XX 22-JUN-2001 (first entry)
DE Amyloid beta-protein fragment peptide SEQ ID NO:976.
XX
XX Protection; endogenous therapeutic peptide; peptidase; conjugation;
KW blood component; modification; succinimidyl; maleimido group; amino;
KW hydroxyl; thiol; hormone; growth factor; neurotransmitter.
XX
XX Homo sapiens.
OS Synthetic.
XX
XX WO200069900-A2.
XX
XX 23-NOV-2000.
XX
XX 17-MAY-2000; 2000WO-US13576.
XX
XX 17-MAY-1999; 99US-0134406.
XX
XX 10-SEP-1999; 99US-0153406.
XX
XX 15-OCT-1999; 99US-0159783.
XX
XX (CONJ-) CONJUCHEM INC.
XX
XX Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;
XX
XX WPI; 2001-112059/12.
XX
XX Modifying and attaching therapeutic peptides to albumin prevents
PT peptidase degradation, useful for increasing length of in vivo activity
PT
PS Disclosure; Page 513; 733pp; English.
XX
XX The present invention describes a modified therapeutic peptide (I)
CC comprising a therapeutically active amino acid region (III) and a
CC reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
CC a less therapeutically active amino acid region (IV), which covalently
CC bonds with amino/hydroxyl/thiol groups on blood components to form a
CC peptide stabilised therapeutic peptide composed of 3-50 amino acids.
CC (I) are useful for modifying therapeutic peptides e.g. hormones, growth
CC factors and neurotransmitters, to protect them from peptidase activity
CC in vivo for the treatment of various disorders. Endogenous therapeutic
CC administration due to rapid degradation by peptidases as they require frequent
CC peptides are not suitable as drug candidates to increase length of activity (half
CC life) and specificity as bonding to large molecules decreases
CC intracellular uptake and interference with physiological processes.
CC AAB90829 to AAB92441 represent peptides which can be used in the
XX exemplification of the present invention.
XX
XX Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
DB 13 HHQKLVFFAE 22

RESULT 31
AAB91816
ID AAB91816 standard; Peptide; 28 AA.
XX
XX AAB91816;
XX
XX 22-JUN-2001 (first entry)
DE Amyloid beta-protein fragment peptide SEQ ID NO:992.
XX
XX Protection; endogenous therapeutic peptide; peptidase; conjugation;
KW blood component; modification; succinimidyl; maleimido group; amino;
KW hydroxyl; thiol; hormone; growth factor; neurotransmitter.
XX
XX Homo sapiens.
OS Synthetic.
XX
XX WO200069900-A2.
XX
XX 23-NOV-2000.
XX
XX 17-MAY-2000; 2000WO-US13576.
XX
XX 17-MAY-1999; 99US-0134406.
XX
XX 10-SEP-1999; 99US-0153406.
XX
XX 15-OCT-1999; 99US-0159783.
XX
XX (CONJ-) CONJUCHEM INC.
XX
XX Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;
XX
XX WPI; 2001-112059/12.
XX
XX Modifying and attaching therapeutic peptides to albumin prevents
PT peptidase degradation, useful for increasing length of in vivo activity
PT
PS Disclosure; Page 519; 733pp; English.
XX
XX The present invention describes a modified therapeutic peptide (I)
CC comprising a therapeutically active amino acid region (III) and a
CC reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
CC a less therapeutically active amino acid region (IV), which covalently
CC bonds with amino/hydroxyl/thiol groups on blood components to form a
CC peptide stabilised therapeutic peptide composed of 3-50 amino acids.
CC (I) are useful for modifying therapeutic peptides e.g. hormones, growth
CC factors and neurotransmitters, to protect them from peptidase activity
CC in vivo for the treatment of various disorders. Endogenous therapeutic
CC administration due to rapid degradation by peptidases as they require frequent
CC peptides are not suitable as drug candidates to increase length of activity (half
CC life) and specificity as bonding to large molecules decreases
CC intracellular uptake and interference with physiological processes.
CC AAB90829 to AAB92441 represent peptides which can be used in the
XX exemplification of the present invention.
XX
XX Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 HHQKLVFFAE 10
DB 13 HHQKLVFFAE 22

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XX	AAB49396;
AC	06-MAR-2001 (first entry)
XX	Human amyloid peptide protein fragment SEQ ID NO: 11.
DT	Human; immunogenic peptide; immune response; monophosphoryl lipid A;
XX	antigen; infection; cancer; amyloid deposition.
DE	Homo sapiens.
XX	WO2000069456-A2.
KW	23-NOV-2000.
KW	12-MAY-2000; 2000WO-US13156.
OS	13-MAY-1999; 99US-0133963.
XX	(AMCY) AMERICAN CYANAMID CO.
XX	Hagen M;
PI	WPI; 2001-024946/03.
XX	Antigenic composition having an antigen (e.g. viral protein) and an
DR	adjuvant, useful for enhancing humoral and cellular immune response in
XX	a host or as a prophylaxis against virus, bacterium, parasite, cancer
PT	cell or allergen -
PT	Disclosure; Page 40; 129pp; English.
XX	The present invention provides an antigenic composition comprising an
CC	lipid A adjuvant. The presence of the adjuvant causes an increased immune
CC	response. The antigen may be from a pathogenic bacterium, fungus, virus,
CC	or parasite, a cancer cell, an allergen or from amyloid peptide protein.
CC	The composition can be used in the prevention and treatment of infection,
CC	cancer and diseases caused by amyloid deposition. It is particularly
CC	useful against HIV, Neisseria gonorrhoeae and respiratory syncytial
CC	virus.
XX	Sequence 28 AA;
SQ	Query Match 100.0%; Score 55; DB 22; Length 28;
	Best Local Similarity 100.0%; Pred. No. 0.00044;
	Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Oy	1 HHQKLVFFAE 10
Dd	
	13 HHQKLVFFAE 22
RESULT 34	
AAB35590	ID AAB35590 standard; peptide; 28 AA.
XX	AC AAB35590;
XX	15-FEB-2001 (first entry)
DT	Human clone B(1-28) amyloid B peptide.
DE	Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
XX	acute cardiovascular disease; therapy.
KW	Homo sapiens.
KW	US6136548-A.
XX	24-OCT-2000.
OS	
XX	
PN	
XX	
PD	
XX	

RESULT 32	
AAB91827	
ID	AAB91827 standard; Peptide; 28 AA.
XX	
AC	AAB91827;
XX	
DT	22-JUN-2001 (first entry)
XX	
XX	Amyloid beta-protein fragment peptide SEQ ID NO:1003.
DE	
XX	
KW	Protection; endogenous therapeutic peptide; peptidase; conjugation;
KW	blood component; modification; succinimidyl; maleimido group; amino;
KW	hydroxyl; thiol; hormone; growth factor; neurotransmitter.
XX	
OS	Homo sapiens.
OS	Synthetic.
XX	
PN	WO20069900-A2.
XX	
PD	23-NOV-2000.
XX	
PF	17-MAY-2000; 2000WO-US13576.
XX	
PR	17-MAY-1999; 99US-0134406.
PR	10-SEP-1999; 99US-0153406.
PR	15-OCT-1999; 99US-0159783.
XX	
PA	(CONJ-) CONJUCHEM INC.
XX	
PI	Bridon DP, Ezrin AM, Milner PG, Holmes DL, Thibaudeau K;
XX	
DR	WPI; 2001-112059/12.
XX	
PT	Modifying and attaching therapeutic peptides to albumin prevents
PT	peptidase degradation, useful for increasing length of in vivo activity
PT	
XX	
PS	Disclosure; Page 523; 733pp; English.
XX	
CC	The present invention describes a modified therapeutic peptide (I)
CC	comprising a therapeutically active amino acid region (III) and a
CC	reactive group (II) (e.g. succinimidyl and maleimido groups) attached to
CC	a less therapeutically active amino acid region (IV), which covalently
CC	bonds with amino/hydroxy/thiol groups on blood components to form a
CC	peptidase stabilised therapeutic peptide composed of 3-50 amino acids.
CC	(I) are useful for modifying therapeutic peptides e.g. hormones, growth
CC	factors and neurotransmitters, to protect them from peptidase activity
CC	in vivo for the treatment of various disorders. Endogenous therapeutic
CC	peptides are not suitable as drug candidates as they require frequent
CC	administration due to rapid degradation by peptidases in the body.
CC	Modifying and attaching therapeutic peptides to albumin prevents or
CC	reduces the action of peptidases to increase length of activity (half
CC	life) and specificity as bonding to large molecules decreases
CC	intracellular uptake and interference with physiological processes.
CC	AAB90829 to AAB92441 represent peptides which can be used in the
CC	exemplification of the present invention.
XX	
SQ	Sequence 28 AA;
	Query Match 100.0%; Score 55; DB 22; Length 28;
	Best Local Similarity 100.0%; Pred. No. 0.00044;
	Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy	1 HHQKLVFAE 10
Db	13 HHQKLVFAE 22
RESULT 33	
AAB49396	
ID	AAB49396 standard; peptide; 28 AA.

```
PF 02-SEP-1999; 99US-0388890.
XX
PR 26-JUL-1996; 96US-0686959.
PR 22-NOV-1994; 94US-0347144.
XX 22-NOV-1995; 95WO-US15007.
PA (RUTF ) UNIV RUTGERS STATE NEW JERSEY.
XX
PI Anderson S;
XX
DR WPI; 2001-030939/04.
XX
PT Identifying mutant tissue-type plasminogen activator (t-PA) for
PT improving thrombolytic therapy or treating vascular hemorrhaging, by
PT determining whether t-PA binds to fibrin but not to a beta amyloid
PT peptide -
XX
PS Example 3; Column 26; 23pp; English.
XX
CC The present invention describes a method for identifying mutant
CC derivatives of tissue-type plasminogen activator, which involves
CC determining whether or not they bind to beta-amyloid peptides and fibrin.
CC Mutants will only bind to the latter. These mutants are useful in
CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
CC and in the treatment of acute cardiovascular disease, which may be caused
CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
XX
SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
DB 13 HHQKLVFFAE 22
RESULT 35
AAB35591
ID AAB35591 standard; peptide; 28 AA.
XX
AC AAB35591;
XX
DT 15-FEB-2001 (first entry)
XX
DE Human clone DIN B(1-28) amyloid B peptide.
XX
KW Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
KW acute cardiovascular disease; therapy.
XX
OS Homo sapiens.
XX
PN US6136548-A.
XX
PD 24-OCT-2000.
XX
PF 02-SEP-1999; 99US-0388890.
XX
PR 26-JUL-1996; 96US-0686959.
PR 22-NOV-1994; 94US-0347144.
XX 22-NOV-1995; 95WO-US15007.
PA (RUTF ) UNIV RUTGERS STATE NEW JERSEY.
XX
PI Anderson S;
XX
DR WPI; 2001-030939/04.
XX
PT Identifying mutant tissue-type plasminogen activator (t-PA) for
PT improving thrombolytic therapy or treating vascular hemorrhaging, by
PT determining whether t-PA binds to fibrin but not to a beta amyloid
PT peptide -
```

```
XX
PS Example 3; Column 26; 23pp; English.
XX
CC The present invention describes a method for identifying mutant
CC derivatives of tissue-type plasminogen activator, which involves
CC determining whether or not they bind to beta-amyloid peptides and fibrin.
CC Mutants will only bind to the latter. These mutants are useful in
CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
CC and in the treatment of acute cardiovascular disease, which may be caused
CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
XX
SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 HHQKLVFFAE 10
DB 13 HHQKLVFFAE 22
RESULT 36
AAB35592
ID AAB35592 standard; peptide; 28 AA.
XX
AC AAB35592;
XX
DT 15-FEB-2001 (first entry)
XX
DE Human clone E3Q B(1-28) amyloid B peptide.
XX
KW Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
KW acute cardiovascular disease; therapy.
XX
OS Homo sapiens.
XX
PN US6136548-A.
XX
PD 24-OCT-2000.
XX
PF 02-SEP-1999; 99US-0388890.
XX
PR 26-JUL-1996; 96US-0686959.
PR 22-NOV-1994; 94US-0347144.
XX 22-NOV-1995; 95WO-US15007.
PA (RUTF ) UNIV RUTGERS STATE NEW JERSEY.
XX
PI Anderson S;
XX
DR WPI; 2001-030939/04.
XX
PT Identifying mutant tissue-type plasminogen activator (t-PA) for
PT improving thrombolytic therapy or treating vascular hemorrhaging, by
PT determining whether t-PA binds to fibrin but not to a beta amyloid
PT peptide -
XX
PS Example 3; Column 26; 23pp; English.
XX
CC The present invention describes a method for identifying mutant
CC derivatives of tissue-type plasminogen activator, which involves
CC determining whether or not they bind to beta-amyloid peptides and fibrin.
CC Mutants will only bind to the latter. These mutants are useful in
CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
CC and in the treatment of acute cardiovascular disease, which may be caused
CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
XX
SQ Sequence 28 AA;
Query Match 100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
```

```

QY      1 HHQKLVFFAE 10
Db      13 HHQKLVFFAE 22

RESULT 37
AAB35593
ID      AAB35593 standard; peptide; 28 AA.
XX
AC      AAB35593;
XX
XX      15-FEB-2001 (first entry)
XX
XX      Human clone R5Q B(1-28) amyloid B peptide.
DE
KW      Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
KW      acute cardiovascular disease; therapy.
XX
OS      Homo sapiens.
XX
PN      US6136548-A.
PD      24-OCT-2000.
XX
PF      02-SEP-1999; 99US-0388890.
XX
PR      26-JUL-1996; 96US-0686959.
PR      22-NOV-1994; 94US-0347144.
PR      22-NOV-1995; 95WO-US15007.
XX
XX      (RUTF ) UNIV RUTGERS STATE NEW JERSEY.
PA
XX
XX      Anderson S;
XX
XX      WPI; 2001-030939/04.
XX
XX      Identifying mutant tissue-type plasminogen activator (t-PA) for
XX      improving thrombolytic therapy or treating vascular hemorrhaging, by
XX      determining whether t-PA binds to fibrin but not to a beta amyloid
XX      peptide.
XX
XX      Example 3; Column 26; 23pp; English.
XX
XX      The present invention describes a method for identifying mutant
XX      derivatives of tissue-type plasminogen activator, which involves
XX      determining whether or not they bind to beta-amyloid peptides and fibrin.
XX      Mutants will only bind to the latter. These mutants are useful in
XX      improved thrombolytic therapies, in the treatment of Alzheimer's disease
XX      and in the treatment of acute cardiovascular disease, which may be caused
XX      by myocardial infarction, stroke, ischaemia and pulmonary embolism.
XX
SQ      Sequence 28 AA;

Query Match      100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY      1 HHQKLVFFAE 10
Db      13 HHQKLVFFAE 22

RESULT 39
AAB35595
ID      AAB35595 standard; peptide; 28 AA.
XX
XX      AAB35595;
AC
XX
XX      15-FEB-2001 (first entry)
XX
XX      Human clone D7Q B(1-28) amyloid B peptide.
XX
XX      Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
XX      acute cardiovascular disease; therapy.
XX
OS      Homo sapiens.
XX
PN      US6136548-A.
PD      24-OCT-2000.
XX
PF      02-SEP-1999; 99US-0388890.
XX
PR      26-JUL-1996; 96US-0686959.
PR      22-NOV-1994; 94US-0347144.
PR      22-NOV-1995; 95WO-US15007.
XX
XX      (RUTF ) UNIV RUTGERS STATE NEW JERSEY.
PA
XX
XX      Anderson S;
XX
XX      WPI; 2001-030939/04.
XX
XX      Identifying mutant tissue-type plasminogen activator (t-PA) for
XX      improving thrombolytic therapy or treating vascular hemorrhaging, by
XX      determining whether t-PA binds to fibrin but not to a beta amyloid
XX      peptide.
XX
XX      Example 3; Column 26; 23pp; English.
XX
XX      The present invention describes a method for identifying mutant
XX      derivatives of tissue-type plasminogen activator, which involves
XX      determining whether or not they bind to beta-amyloid peptides and fibrin.
XX      Mutants will only bind to the latter. These mutants are useful in
XX      improved thrombolytic therapies, in the treatment of Alzheimer's disease
XX      and in the treatment of acute cardiovascular disease, which may be caused
XX      by myocardial infarction, stroke, ischaemia and pulmonary embolism.
XX
SQ      Sequence 28 AA;

Query Match      100.0%; Score 55; DB 22; Length 28;
Best Local Similarity 100.0%; Pred. No. 0.00044;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY      1 HHQKLVFFAE 10
Db      13 HHQKLVFFAE 22

RESULT 38
AAB35594
ID      AAB35594 standard; peptide; 28 AA.
XX
XX      AAB35594;
AC
XX
XX      15-FEB-2001 (first entry)
XX
XX      Human clone H6Q B(1-28) amyloid B peptide.
DE
XX

```

XX Anderson S;
 XX WPI; 2001-030939/04.
 XX Identifying mutant tissue-type plasminogen activator (t-PA) for
 PT improving thrombolytic therapy or treating vascular hemorrhaging, by
 PT determining whether t-PA binds to fibrin but not to a beta amyloid
 PT peptide -
 XX
 XX Example 3; Column 26; 23pp; English.
 XX
 CC The present invention describes a method for identifying mutant
 CC derivatives of tissue-type plasminogen activator, which involves
 CC determining whether or not they bind to beta-amyloid peptides and fibrin.
 CC Mutants will only bind to the latter. These mutants are useful in
 CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
 CC and in the treatment of acute cardiovascular disease, which may be caused
 CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
 XX
 XX Sequence 28 AA;
 SQ
 Query Match 100.0%; Score 55; DB 22; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 13 HHQKLVFFAE 22
 |||||
 RESULT 40
 AAB35596
 ID AAB35596 standard; peptide; 28 AA.
 XX
 AC AAB35596;
 XX
 DT 15-FEB-2001 (first entry)
 XX
 DE Human clone E11Q B(1-28) amyloid B peptide.
 XX
 KW Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
 KW acute cardiovascular disease; therapy.
 XX
 OS Homo sapiens.
 XX
 PN US6136548-A.
 XX
 PD 24-OCT-2000.
 XX
 PF 02-SEP-1999; 99US-0388890.
 XX
 PR 26-JUL-1996; 96US-0686959.
 PR 22-NOV-1994; 94US-0347144.
 PR 22-NOV-1995; 95WO-US15007.
 XX
 PA (RUTF) UNIV RUTGERS STATE NEW JERSEY.
 XX
 PI Anderson S;
 XX
 DR WPI; 2001-030939/04.
 XX
 PN Identifying mutant tissue-type plasminogen activator (t-PA) for
 PT improving thrombolytic therapy or treating vascular hemorrhaging, by
 PT determining whether t-PA binds to fibrin but not to a beta amyloid
 PT peptide -
 XX
 XX Example 3; Column 26; 23pp; English.
 XX
 CC The present invention describes a method for identifying mutant
 CC derivatives of tissue-type plasminogen activator, which involves
 CC determining whether or not they bind to beta-amyloid peptides and fibrin.
 CC Mutants will only bind to the latter. These mutants are useful in

CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
 CC and in the treatment of acute cardiovascular disease, which may be caused
 CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
 XX
 XX Sequence 28 AA;
 SQ
 Query Match 100.0%; Score 55; DB 22; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 13 HHQKLVFFAE 22
 |||||
 RESULT 41
 AAB36201
 ID AAB36201 standard; peptide; 28 AA.
 XX
 AC AAB36201;
 XX
 DT 15-FEB-2001 (first entry)
 XX
 DE Human clone D23Q B(1-28) amyloid B peptide.
 XX
 KW Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
 KW acute cardiovascular disease; therapy.
 XX
 OS Homo sapiens.
 XX
 PN US6136548-A.
 XX
 PD 24-OCT-2000.
 XX
 PF 02-SEP-1999; 99US-0388890.
 XX
 PR 26-JUL-1996; 96US-0686959.
 PR 22-NOV-1994; 94US-0347144.
 PR 22-NOV-1995; 95WO-US15007.
 XX
 PA (RUTF) UNIV RUTGERS STATE NEW JERSEY.
 XX
 PI Anderson S;
 XX
 DR WPI; 2001-030939/04.
 XX
 PN Identifying mutant tissue-type plasminogen activator (t-PA) for
 PT improving thrombolytic therapy or treating vascular hemorrhaging, by
 PT determining whether t-PA binds to fibrin but not to a beta amyloid
 PT peptide -
 XX
 XX Example 3; Column 26; 23pp; English.
 XX
 CC The present invention describes a method for identifying mutant
 CC derivatives of tissue-type plasminogen activator, which involves
 CC determining whether or not they bind to beta-amyloid peptides and fibrin.
 CC Mutants will only bind to the latter. These mutants are useful in
 CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
 CC and in the treatment of acute cardiovascular disease, which may be caused
 CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
 XX
 XX Sequence 28 AA;
 SQ
 Query Match 100.0%; Score 55; DB 22; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 Db 13 HHQKLVFFAE 22
 |||||
 RESULT 42

AAB36202
 ID AAB36202 standard; peptide; 28 AA.
 XX
 AC AAB36202;
 XX
 DT 15-FEB-2001 (first entry)
 XX
 XX Human clone K28Q B(1-28) amyloid B peptide.
 DE
 XX Beta-amyloid; amyloid deposit; Alzheimer's disease; thrombolytic therapy;
 KW acute cardiovascular disease; therapy.
 XX
 XX Homo sapiens.
 OS
 XX US6136548-A.
 PN
 XX 24-OCT-2000.
 PD
 XX 02-SEP-1999; 99US-0388890.
 XX
 XX 26-JUL-1996; 96US-0686959.
 PR
 XX 22-NOV-1994; 94US-0347144.
 PR
 XX 22-NOV-1995; 95WO-US15007.
 XX
 XX (RUTF) UNIV RUTGERS STATE NEW JERSEY.
 PA
 XX Anderson S;
 PI
 XX WPI; 2001-030939/04.
 DR
 XX Identifying mutant tissue-type plasminogen activator (t-PA) for
 PT improving thrombolytic therapy or treating vascular hemorrhaging, by
 PT determining whether t-PA binds to fibrin but not to a beta amyloid
 PT peptide -
 XX
 PS Example 3; Column 26; 23pp; English.
 XX
 CC The present invention describes a method for identifying mutant
 CC derivatives of tissue-type plasminogen activator, which involves
 CC determining whether or not they bind to beta-amyloid peptides and fibrin.
 CC Mutants will only bind to the latter. These mutants are useful in
 CC improved thrombolytic therapies, in the treatment of Alzheimer's disease
 CC and in the treatment of acute cardiovascular disease, which may be caused
 CC by myocardial infarction, stroke, ischaemia and pulmonary embolism.
 XX
 SQ Sequence 28 AA;
 Query Match 100.0%; Score 55; DB 22; Length 28;
 Best Local Similarity 100.0%; Pred. No. 0.00044;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 DB 13 HHQKLVFFAE 22
 RESULT 43
 AAW81468
 ID AAW81468 standard; peptide; 30 AA.
 XX
 AC AAW81468;
 XX
 DT 28-JAN-1999 (first entry)
 XX
 DE Synthetic amyloid beta (Abeta) peptide 3 (residues 1-30).
 XX
 KW Amyloid beta; Abeta; deoxygenated solvent; evaporative deposition;
 KW research; neurotoxicity; free-radical; glutamine synthetase.
 XX
 OS Synthetic.
 XX
 PN US5840838-A.
 XX

PD 24-NOV-1998.
 XX
 PF 29-FEB-1996; 36US-0609090.
 XX
 PR 29-FEB-1996; 36US-0609090.
 XX
 PA (KENT) UNIV KENTUCKY RES FOUND.
 XX
 XX Aksenov M, Butterfield DA, Carney JM, Hensley K;
 PI WPI; 1999-034120/03.
 DR
 XX Process for treating synthetic amyloid beta peptides - by organic
 PT solvent treatment, useful for studying neurotoxicity
 PT
 XX Claim 5; Columns 9-10; 14pp; English.
 PS
 XX Sequences AAW81466 to AAW81476 represent synthetic amyloid beta (Abeta)
 CC peptides. The invention provides a process for treating a synthetic
 CC Abeta peptide that comprises dissolving the peptide in a deoxygenated
 CC solvent selected from trifluoroethanol, hexafluorocyclohexane, dimethyl
 CC sulphoxide, morpholinopropanesulphonic acid, dimethylformamide and
 CC acetonitrile to a concentration of 0.01-10 mg/ml, incubating the
 CC solution at 20-65 deg. C for 0.5-4 hour, and removing the solvent by
 CC //evaporative deposition// in 5-10 minutes. Synthetic amyloid beta
 CC peptides are useful as research tools for studying neurotoxicity
 CC resulting from Abeta peptide -enhanced free-radical production. The
 CC treatment increases the activity of the synthetic Abeta peptides in tests
 CC to determine free-radical generating capacity and glutamine synthetase
 CC inactivation.
 XX
 SQ Sequence 30 AA;
 Query Match 100.0%; Score 55; DB 20; Length 30;
 Best Local Similarity 100.0%; Pred. No. 0.00047;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHQKLVFFAE 10
 DB 13 HHQKLVFFAE 22
 RESULT 44
 AAB84430
 ID AAB84430 standard; peptide; 32 AA.
 XX
 AC AAB84430;
 XX
 DT 22-AUG-2001 (first entry)
 XX
 DE Partial sequence of a human beta-amyloid precursor protein.
 XX
 KW Beta-amyloid precursor protein; APP; chimeric peptide; B cell epitope;
 KW vaccine.
 XX
 OS Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Modified-site 1
 FT /note= "pyroglutamate"
 XX
 PN WO200142306-A2.
 XX
 PD 14-JUN-2001.
 XX
 PF 08-DEC-2000; 2000WO-US33203.
 XX
 PR 08-DEC-1999; 99US-0169687.
 XX
 PA (MIND-) MINDSET BIOPHARMACEUTICALS USA INC.
 XX
 PI Chain B;
 XX

DR WPI; 2001-381648/40.
 XX Novel chimeric peptide containing N- or C-terminal end-specific B cell
 PT epitope from naturally occurring internal peptide cleavage product
 PT (such as beta amyloid peptide) of a precursor protein, joined to T cell
 PT epitope
 XX
 PS Claim 3; Page 42-43; 47pp; English.
 XX
 CC The present sequence represents a partial sequence of a human
 CC beta-amyloid precursor protein (APP). The peptide is used to create
 CC chimeric peptides of the invention. The chimeric peptides contain a N-
 CC or C-terminal end-specific B cell epitope from a naturally occurring
 CC internal peptide cleavage product of a precursor or mature protein, as
 CC a free N- or C-terminus, joined to a T cell epitope, with or without a
 CC spacer amino acid residue. Chimeric peptides comprising betaAPP peptides
 CC slow down, reduce or prevent the accumulation of amyloid beta peptide in
 CC the extracellular space, interstitial fluid and cerebrospinal fluid of
 CC the brain, and aggregation into senile amyloid deposits or plaques. They
 CC also block the interaction of amyloid beta peptides with other molecules
 CC that contribute the neurotoxicity of amyloid beta. The chimeric peptides
 CC are useful for immunizing humans against the free N- or C-terminus of
 CC an internal self peptide cleavage product (e.g. APP peptide) derived from
 CC a precursor protein or a mature protein. The internal peptide cleavage
 CC product is the self molecule of the mammal.
 XX
 SQ Sequence 32 AA;
 Query Match 100.0%; Score 55; DB 22; Length 32;
 Best Local Similarity 100.0%; Pred. No. 0.0005;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHOKLVFFAE 10
 Db 3 HHOKLVFFAE 12
 RESULT 45
 AAW81469
 ID AAW81469 standard; peptide; 33 AA.
 XX
 AC AAW81469;
 XX
 DT 28-JAN-1999 (first entry)
 XX
 DE Synthetic amyloid beta (Abeta) peptide 4 (residues 1-33).
 XX
 KW Amyloid beta; Abeta; deoxygenated solvent; evaporative deposition;
 KW research; neurotoxicity; free-radical; glutamine synthetase.
 XX
 OS Synthetic.
 XX
 PN US5840838-A.
 XX
 PD 24-NOV-1998.
 XX
 PF 29-FEB-1996; 96US-0609090.
 XX
 PR 29-FEB-1996; 96US-0609090.
 XX
 PS (KENT) UNIV KENTUCKY RES FOUND.
 PA
 PI Aksenov M, Butterfield DA, Carney JM, Hensley K;
 XX
 XX WPI; 1999-034120/03.
 DR
 XX Process for treating synthetic amyloid beta peptides - by organic
 PT solvent treatment, useful for studying neurotoxicity
 XX
 PS Claim 5; Columns 9-10; 14pp; English.
 XX
 CC Sequences AAW81466 to AAW81476 represent synthetic amyloid beta (Abeta)
 CC peptides. The invention provides a process for treating a synthetic

CC Abeta peptide that comprises dissolving the peptide in a deoxygenated
 CC solvent selected from trifluoroethanol, hexafluorocyclohexane, dimethyl
 CC sulphoxide, morpholinopropanesulphonic acid, dimethylformamide and
 CC acetonitrile to a concentration of 0.01-10 mg/ml, incubating the
 CC solution at 20-65 deg. C for 0.5-4 hour, and removing the solvent by
 CC "evaporative deposition" in 5-10 minutes. Synthetic amyloid beta
 CC peptides are useful as research tools for studying neurotoxicity
 CC resulting from Abeta peptide-enhanced free-radical production. The
 CC treatment increases the activity of the synthetic Abeta peptides in tests
 CC to determine free-radical generating capacity and glutamine synthetase
 CC inactivation.
 XX
 SQ Sequence 33 AA;
 Query Match 100.0%; Score 55; DB 20; Length 33;
 Best Local Similarity 100.0%; Pred. No. 0.00052;
 Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 HHOKLVFFAE 10
 Db 13 HHOKLVFFAE 22
 RESULT 46
 AAW47228
 ID AAW47228 standard; peptide; 35 AA.
 XX
 AC AAW47228;
 XX
 DT 22-MAY-1998 (first entry)
 XX
 DE Beta-amyloid peptide residues 1-35.
 XX
 KW Screening assay; beta-amyloid peptide; treatment;
 KW amyloidosis disease; Alzheimer's disease.
 XX
 OS Homo sapiens.
 XX
 PN US5721106-A.
 XX
 PD 24-FEB-1998.
 XX
 PF 12-SEP-1994; 94US-0304585.
 XX
 PR 12-SEP-1994; 94US-0304585.
 PR 13-AUG-1991; 91US-0744767.
 XX
 PA (HARD) HARVARD COLLEGE.
 PA (MINU) UNIV MINNESOTA.
 XX
 PI Magglo JE, Mantyh PW;
 XX
 DR WPI; 1998-168404/15.
 XX
 PT New in vitro screening assay for Alzheimer's disease drugs -
 PT comprises assessing binding of labelled beta-amyloid peptide to silk
 PT sample
 XX
 PS Claim 8; Columns 31-32; 36pp; English.
 XX
 CC The present sequence was used in the development of a novel in
 CC vitro screening assay for agents capable of affecting the
 CC deposition of beta-amyloid peptide (BAP) on tissue. The method
 CC comprises contacting a silk sample with labelled BAP, optionally
 CC in the presence of a test agent, detecting the amount of label
 CC bound to the silk and assessing the effect of the agent on the
 CC deposition of BAP. Agents that inhibit binding of BAP to silk are
 CC potentially useful for treating amyloidosis diseases, especially
 CC Alzheimer's disease.
 XX
 SQ Sequence 35 AA;
 Query Match 100.0%; Score 55; DB 19; Length 35;

```
Best Local Similarity 100.0%; Pred. No. 0.00056;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVEFAE 10
DB 13 HHQKLVEFAE 22

RESULT 47
AAW89357
ID AAW89357 standard; peptide; 35 AA.
XX
AC AAW89357;
XX
DT 02-MAR-1999 (first entry)
XX
DE Beta-amyloid peptide derivative A-beta-6-40.
XX
KW Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
familial amyloid polyneuropathy; bovine spongiform encephalopathy;
KW Creutzfeldt-Jakob disease; BAP.
XX
OS Homo sapiens.
OS Synthetic.
XX
PN US5854204-A.
XX
PD 29-DEC-1998.
XX
PF 14-MAR-1996; 96US-0612785.
XX
PR 14-MAR-1996; 96US-0612785.
PR 14-MAR-1995; 95US-0404831.
PR 07-JUN-1995; 95US-0475579.
PR 27-OCT-1995; 95US-0548998.
XX
PA (PRAE-) PRAECIS PHARM INC.
XX
PI Benjamin H, Chin J, Findeis MA, Garnick MB, Gefter ML;
PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
PI Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;
XX
DR WPI; 1999-094964/08.
XX
PT New peptide(s) derived from beta-amyloid peptide that inhibit
PT amyloid aggregation - and neurotoxicity, specifically for treatment
PT and prevention of Alzheimer's disease
XX
PS Claim 5; Column 81-82; 52pp; English.
XX
CC The present invention describes beta-amyloid peptide (BAP) derivatives.
CC The BAP derivatives inhibit aggregation of amyloidogenic proteins and
CC peptides, specifically BAP, and their neurotoxicity, so are useful for
CC treating and preventing any disease involving amyloidosis, specifically
CC Alzheimer's disease but also Down's syndrome, familial amyloid
CC polyneuropathy or cardiomyopathy. bovine spongiform encephalopathy and
CC Creutzfeldt-Jakob disease. The BAP derivatives are also used to diagnose
CC these diseases, in vitro or in vivo, by detecting binding of BAP to
CC labelled BAP derivatives. Some BAP derivatives inhibit BAP aggregation
CC even when BAP is present in molar excess. The present sequence
CC represents a BAP derivative.
XX
SQ Sequence 35 AA;
Query Match 100.0%; Score 55; DB 20; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00056;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVEFAE 10
DB 8 HHQKLVEFAE 17

RESULT 48
AAW89359
ID AAW89359 standard; peptide; 35 AA.
XX
AC AAW89359;
XX
DT 02-MAR-1999 (first entry)
XX
DE Beta-amyloid peptide derivative A-beta-1-25,31-40 (Delta26-30).
XX
KW Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
familial amyloid polyneuropathy; bovine spongiform encephalopathy;
KW Creutzfeldt-Jakob disease; BAP.
XX
OS Homo sapiens.
OS Synthetic.
XX
PN US5854204-A.
XX
PD 29-DEC-1998.
XX
PF 14-MAR-1996; 96US-0612785.
XX
PR 14-MAR-1996; 96US-0612785.
PR 14-MAR-1995; 95US-0404831.
PR 07-JUN-1995; 95US-0475579.
PR 27-OCT-1995; 95US-0548998.
XX
PA (PRAE-) PRAECIS PHARM INC.
XX
PI Benjamin H, Chin J, Findeis MA, Garnick MB, Gefter ML;
PI Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
PI Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;
XX
DR WPI; 1999-094964/08.
XX
PT New peptide(s) derived from beta-amyloid peptide that inhibit
PT amyloid aggregation - and neurotoxicity, specifically for treatment
PT and prevention of Alzheimer's disease
XX
PS Claim 7; Column 81-82; 52pp; English.
XX
CC The present invention describes beta-amyloid peptide (BAP) derivatives.
CC The BAP derivatives inhibit aggregation of amyloidogenic proteins and
CC peptides, specifically BAP, and their neurotoxicity, so are useful for
CC treating and preventing any disease involving amyloidosis, specifically
CC Alzheimer's disease but also Down's syndrome, familial amyloid
CC polyneuropathy or cardiomyopathy. bovine spongiform encephalopathy and
CC Creutzfeldt-Jakob disease. The BAP derivatives are also used to diagnose
CC these diseases, in vitro or in vivo, by detecting binding of BAP to
CC labelled BAP derivatives. Some BAP derivatives inhibit BAP aggregation
CC even when BAP is present in molar excess. The present sequence
CC represents a BAP derivative.
XX
SQ Sequence 35 AA;
Query Match 100.0%; Score 55; DB 20; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00056;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHQKLVEFAE 10
DB 13 HHQKLVEFAE 22

RESULT 49
AAW89361
ID AAW89361 standard; peptide; 35 AA.
XX
AC AAW89361;
XX
```

DT XX 02-MAR-1999 (first entry)
DE XX Beta-amyloid peptide derivative A-beta-1-5,11-40 (Delta6-10).
XX
KW Human; beta-amyloid peptide; Alzheimer's disease; amyloidogenic protein;
KW aggregation; neurotoxicity; amyloidosis; Down's syndrome; cardiomyopathy;
KW familial amyloid polyneuropathy; bovine spongiform encephalopathy;
KW Creutzfeldt-Jakob disease; BAP.
XX
OS Homo sapiens.
OS Synthetic.
XX
XX US5854204-A.
XX 29-DEC-1998.
XX
XX 14-MAR-1996; 96US-0612785.
XX
XX 14-MAR-1996; 96US-0612785.
XX 14-MAR-1995; 95US-0404831.
XX 07-JUN-1995; 95US-0475579.
XX 27-OCT-1995; 95US-0548998.
XX
XX (PRAE-) PRAECIS PHARM INC.
XX
XX Benjamin H, Chin J, Findeis MA, Garnick MB, Geffer ML;
XX Hundal A, Kasman L, Kelley M, Kubasek W, Lee J;
XX Molineaux S, Musso G, Reed M, Signer ER, Wakefield J;
XX WPI; 1999-094964/08.
XX
XX New peptide(s) derived from beta-amyloid peptide that inhibit
XX amyloid aggregation - and neurotoxicity, specifically for treatment
XX and prevention of Alzheimer's disease
XX
XX Claim 9; Column 83-84; 52pp; English.
XX
XX The present invention describes beta-amyloid peptide (BAP) derivatives.
XX The BAP derivatives inhibit aggregation of amyloidogenic proteins and
XX peptides, specifically BAP, and their neurotoxicity, so are useful for
XX treating and preventing any disease involving amyloidosis, specifically
XX Alzheimer's disease but also Down's syndrome, familial amyloid
XX polyneuropathy or cardiomyopathy, bovine spongiform encephalopathy and
XX Creutzfeldt-Jakob disease. The BAP derivatives are also used to diagnose
XX these diseases, in vitro or in vivo, by detecting binding of BAP to
XX labelled BAP derivatives. Some BAP derivatives inhibit BAP aggregation
XX even when BAP is present in molar excess. The present sequence
XX represents a BAP derivative.
XX
SQ Sequence 35 AA;

Query Match 100.0%; Score 55; DB 20; Length 35;
Best Local Similarity 100.0%; Pred. No. 0.00056;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 8 HHOKLVFFAE 17

RESULT 50
AAW81471
ID AAW81471 standard; peptide; 36 AA.
XX
XX AAW81471;
XX
XX 28-JAN-1999 (first entry)
XX
XX Synthetic amyloid beta (Abeta) peptide 6 (residues 1-36).
XX
KW Amyloid beta; Abeta; deoxygenated solvent; evaporative deposition;
KW research; neurotoxicity; free-radical; glutamine synthetase.
XX
XX

OS Synthetic.
XX US5840838-A.
XX
XX 24-NOV-1998.
XX
XX 29-FEB-1996; 96US-0609090.
XX
XX 29-FEB-1996; 96US-0609090.
XX
XX (KENT) UNIV KENTUCKY RES FOUND.
XX
XX Aksenov M, Butterfield DA, Carney JM, Hensley K;
XX WPI; 1999-034120/03.
XX
XX Process for treating synthetic amyloid beta peptides - by organic
XX solvent treatment, useful for studying neurotoxicity
XX
XX Claim 5; Columns 11-12; 14pp; English.
XX
XX Sequences AAW81466 to AAW81476 represent synthetic amyloid beta (Abeta)
XX peptides. The invention provides a process for treating a synthetic
XX Abeta peptide that comprises dissolving the peptide in a deoxygenated
XX solvent selected from trifluoroethanol, hexafluorocyclohexane, dimethyl
XX sulphoxide, morpholinopropanesulphonic acid, dimethylformamide and
XX acetonitrile to a concentration of 0.01-10 mg/ml, incubating the
XX solution at 20-65 deg C for 0.5-4 hour, and removing the solvent by
XX "evaporative deposition," in 5-10 minutes. Synthetic amyloid beta
XX peptides are useful as research tools for studying neurotoxicity
XX resulting from Abeta peptide -enhanced free-radical production. The
XX treatment increases the activity of the synthetic Abeta peptides in tests
XX to determine free-radical generating capacity and glutamine synthetase
XX inactivation.
XX
SQ Sequence 36 AA;

Query Match 100.0%; Score 55; DB 20; Length 36;
Best Local Similarity 100.0%; Pred. No. 0.00057;
Matches 10; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 HHOKLVFFAE 10
Db 13 HHOKLVFFAE 22

Search completed: October 29, 2002, 09:24:07
Job time : 33 secs